



# STIC Search Report

## Biotech-Chem Library

STIC Database Tracking Number: 133526

**TO:** Ralph J Gitomer  
**Location:** 3d65 / 3e71  
**Art Unit:** 1651  
**Thursday, September 30, 2004**  
  
**Case Serial Number:** 10/000437

**From:** Noble Jarrell  
**Location:** Biotech-Chem Library  
**Rem 1B71**  
**Phone:** 272-2556  
  
**Noble.jarrell@uspto.gov**

### Search Notes

=> d his

(FILE 'HOME' ENTERED AT 09:57:54 ON 30 SEP 2004)

FILE 'HCAPLUS' ENTERED AT 09:58:04 ON 30 SEP 2004  
L1 1 US20020086342/PN

FILE 'REGISTRY' ENTERED AT 09:58:22 ON 30 SEP 2004

FILE 'HCAPLUS' ENTERED AT 09:58:26 ON 30 SEP 2004  
L2 TRA L1 1- RN : 11 TERMS

FILE 'REGISTRY' ENTERED AT 09:58:27 ON 30 SEP 2004  
L3 11 SEA L2

FILE 'WPIX' ENTERED AT 09:58:30 ON 30 SEP 2004  
L4 1 US20020086342/PN

=> b hcap

FILE 'HCAPLUS' ENTERED AT 09:58:55 ON 30 SEP 2004  
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FILE COVERS 1907 - 30 Sep 2004 VOL 141 ISS 14  
FILE LAST UPDATED: 29 Sep 2004 (20040929/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

=> d all 11

L1 ANSWER 1 OF 1 HCAPLUS COPYRIGHT 2004 ACS on STN  
AN 2002:331864 HCAPLUS  
DN 136:337033  
ED Entered STN: 03 May 2002  
TI Drug screening for inhibitors of mouse protein tyrosine phosphatase PTPbr7 and their use in regulating nerve growth factor  
IN Schaeffer, Eric  
PA Pfizer Products Inc., USA  
SO Eur. Pat. Appl., 14 pp.  
CODEN: EPXXDW

DT Patent

LA English

IC ICM C12Q001-42

CC 7-1 (Enzymes)

Section cross-reference(s) : 1, 13

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	EP 1201764	A2	20020502	EP 2001-124850	20011018
	EP 1201764	A3	20040107		
	R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR				
	US 2002086342	A1	20020704	US 2001-437	20011031 <-
	JP 2002355066	A2	20021210	JP 2001-334245	20011031
PRAI	US 2000-244539P	P	20001031		

CLASS

PATENT NO.	CLASS	PATENT FAMILY CLASSIFICATION CODES
EP 1201764	ICM	C12Q001-42
EP 1201764	ECLA	C12Q001/42

AB The present invention relates to novel assays for the identification of agents that inhibit the catalytic activity of mouse PTPbr7. Substrates

for dephosphorylation include phosphorylated mitogen-activating kinase and small peptides contain phosphotyrosine residues. An assay for identifying inhibitors of PTPbr7 includes an assay buffer containing 50 mM TRIS, 0.15 M NaCl, 5 mM DTT, 0.1% BSA at pH 7.4 and a volume of 25.μL. The assay is terminated by adding malachite green dye, ammonium molybdate, and Tween-20 with incubation period of 15 min. Thereafter, the optical d. of free inorg. phosphate is spectrophotometrically measured at 620 nm and compared with a set of stds., containing varied amts. of inorg. phosphate. The invention also provides pharmaceutical compns. comprising such agents identified using the assays of the invention. The invention further provides methods of treatment comprising administering such pharmaceutical compns.

ST drug screening inhibitor PTPbr7; protein tyrosine phosphatase br7 human regulation nerve growth factor; fusion protein GST histidine tag PTPbr7

IT Signal transduction, biological  
(PTPbr7 as neg. regulator of nerve growth factor; drug screening for inhibitors of human protein tyrosine phosphatase PTPbr7 and their use in regulating nerve growth factor)

IT Chimeric gene  
RL: BSU (Biological study, unclassified); BUU (Biological use, unclassified); BIOL (Biological study); USES (Uses)  
(PTPbr7 catalytic domain and GST encoded by; drug screening for inhibitors of human protein tyrosine phosphatase PTPbr7 and their use in regulating nerve growth factor)

IT Chimeric gene  
RL: BSU (Biological study, unclassified); BUU (Biological use, unclassified); BIOL (Biological study); USES (Uses)  
(PTPbr7 catalytic domain and histidine (6) tag encoded by; drug screening for inhibitors of human protein tyrosine phosphatase PTPbr7 and their use in regulating nerve growth factor)

IT Axon  
(PTPbr7 inhibitor in stimulating growth of; drug screening for inhibitors of human protein tyrosine phosphatase PTPbr7 and their use in regulating nerve growth factor)

IT Brain  
(PTPbr7 mRNA in; drug screening for inhibitors of human protein tyrosine phosphatase PTPbr7 and their use in regulating nerve growth factor)

IT Drug screening  
Mus  
(drug screening for inhibitors of human protein tyrosine phosphatase PTPbr7 and their use in regulating nerve growth factor)

IT Dephosphorylation, biological  
(of phospho-MAPK or peptides containing phosphotyrosines by PTPbr7, inhibitors of; drug screening for inhibitors of human protein tyrosine phosphatase PTPbr7 and their use in regulating nerve growth factor)

IT Peptides, biological studies  
RL: BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
(phosphotyrosine, PTPbr7 in dephosphorylation of; drug screening for inhibitors of human protein tyrosine phosphatase PTPbr7 and their use in regulating nerve growth factor)

IT 9061-61-4, Nerve growth factor  
RL: BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
(PTPbr7 as neg. regulator of; drug screening for inhibitors of human protein tyrosine phosphatase PTPbr7 and their use in regulating nerve growth factor)

IT 142243-02-5, Mitogen activated protein kinase  
RL: BSU (Biological study, unclassified); BIOL (Biological study)  
(PTPbr7 in dephosphorylation of; drug screening for inhibitors of human protein tyrosine phosphatase PTPbr7 and their use in regulating nerve growth factor)

IT 373389-66-3, PTPBR7  
RL: ADV (Adverse effect, including toxicity); BSU (Biological study, unclassified); BIOL (Biological study)  
(PTPbr7, of mouse; drug screening for inhibitors of human protein tyrosine phosphatase PTPbr7 and their use in regulating nerve growth factor)

IT 148851-08-5 416848-49-2  
RL: BSU (Biological study, unclassified); BUU (Biological use, unclassified); BIOL (Biological study); USES (Uses)  
(amino acid sequence for PTPbr7 peptide substrate; drug screening for inhibitors of human protein tyrosine phosphatase PTPbr7 and their use in regulating nerve growth factor)

IT 13721-39-6, Sodium orthovanadate

RL: BUU (Biological use, unclassified); BIOL (Biological study); USES (Uses)  
 (as non-specific tyrosine phosphate inhibitor in drug screening assay;  
 drug screening for inhibitors of human protein tyrosine phosphatase  
 PTPbr7 and their use in regulating nerve growth factor)

IT 754-02-9  
 RL: BUU (Biological use, unclassified); BIOL (Biological study); USES (Uses)  
 (as substrate for PTPbr7 dephosphorylation in drug screening assay;  
 drug screening for inhibitors of human protein tyrosine phosphatase  
 PTPbr7 and their use in regulating nerve growth factor)

IT 50812-37-8D, Glutathione-S-transferase, PTPbr7 catalytic domain fusion product with  
 RL: BUU (Biological use, unclassified); BIOL (Biological study); USES (Uses)  
 (drug screening for inhibitors of human protein tyrosine phosphatase  
 PTPbr7 and their use in regulating nerve growth factor)

IT 71-00-1D, L-Histidine, PTPbr7 catalytic domain fusion product with  
 RL: BUU (Biological use, unclassified); BIOL (Biological study); USES (Uses)  
 (tag; drug screening for inhibitors of human protein tyrosine phosphatase PTPbr7 and their use in regulating nerve growth factor)

IT 416933-31-8 416933-32-9  
 RL: PRP (Properties)  
 (unclaimed sequence; drug screening for inhibitors of mouse protein tyrosine phosphatase PTPbr7 and their use in regulating nerve growth factor)

=> b reg  
 FILE 'REGISTRY' ENTERED AT 09:59:01 ON 30 SEP 2004  
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Property values tagged with IC are from the ZIC/VINITI data file  
 provided by InfoChem.

STRUCTURE FILE UPDATES: 29 SEP 2004 HIGHEST RN 754169-63-6  
 DICTIONARY FILE UPDATES: 29 SEP 2004 HIGHEST RN 754169-63-6

TSCA INFORMATION NOW CURRENT THROUGH MAY 21, 2004

Please note that search-term pricing does apply when  
 conducting SmartSELECT searches.

Crossover limits have been increased. See HELP CROSSOVER for details.

Experimental and calculated property data are now available. For more  
 information enter HELP PROP at an arrow prompt in the file or refer  
 to the file summary sheet on the web at:  
<http://www.cas.org/ONLINE/DBSS/registryss.html>

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L3 ANSWER 1 OF 11 REGISTRY COPYRIGHT 2004 ACS on STN  
 RN 416933-32-9 REGISTRY  
 CN DNA, d(G-A-C-A-C-T-C-T-G-G-T-A-A-A-T-C-T-T-C-G-G) (9CI) (CA INDEX NAME)  
 OTHER NAMES:  
 CN 4: PN: EP1201764 PAGE: 11 unclaimed DNA  
 FS NUCLEIC ACID SEQUENCE  
 MF Unspecified  
 CI MAN  
 SR CA  
 LC STN Files: CA, CAPLUS, TOXCENTER, USPATFULL  
 DT.CA CAplus document type: Patent  
 RL.P Roles from patents: PRP (Properties)

\*\*RELATED SEQUENCES AVAILABLE WITH SEQLINK\*\*

\*\*\* STRUCTURE DIAGRAM IS NOT AVAILABLE \*\*\*  
 \*\*\* USE 'SQD' OR 'SQIDE' FORMATS TO DISPLAY SEQUENCE \*\*\*  
 1 REFERENCES IN FILE CA (1907 TO DATE)  
 1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

L3 ANSWER 2 OF 11 REGISTRY COPYRIGHT 2004 ACS on STN

RN 416933-31-8 REGISTRY  
 CN DNA, d(C-T-T-A-G-C-C-T-A-G-A-G-T-G-G-G-T-T-G-G-C-G-G-C) (9CI) (CA INDEX  
 NAME)  
 OTHER NAMES:  
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 FS NUCLEIC ACID SEQUENCE  
 MF Unspecified  
 CI MAN  
 SR CA  
 LC STN Files: CA, CAPLUS, TOXCENTER, USPATFULL  
 DT.CA CAplus document type: Patent  
 RL.P Roles from patents: PRP (Properties)

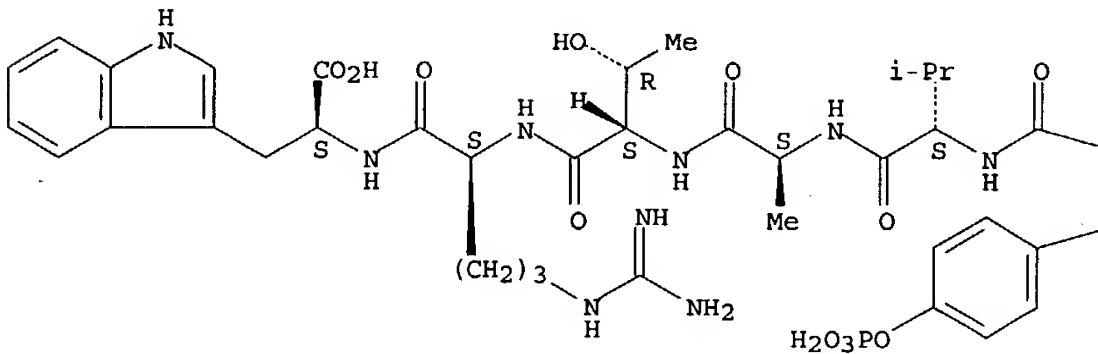
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 1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

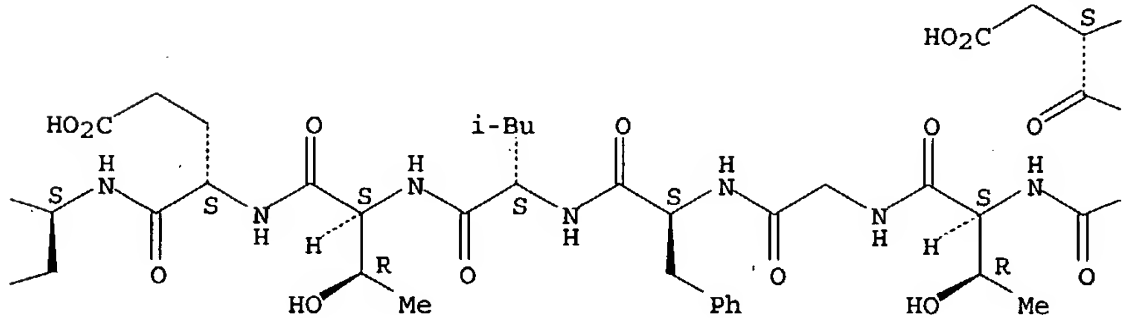
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 RN 416848-49-2 REGISTRY  
 CN L-Tryptophan, L-.alpha.-aspartyl-L-histidyl-L-threonylglycyl-L-  
 phenylalanyl-L-leucyl-L-threonyl-L-.alpha.-glutamyl-O-phosphono-L-tyrosyl-  
 L-valyl-L-alanyl-L-threonyl-L-arginyl- (9CI) (CA INDEX NAME)  
 OTHER NAMES:  
 CN 2: PN: EP1201764 PAGE: 13 claimed sequence  
 FS PROTEIN SEQUENCE; STEREOSEARCH  
 MF C78 H111 N20 O26 P  
 SR CA  
 LC STN Files: CA, CAPLUS, TOXCENTER, USPATFULL  
 DT.CA CAplus document type: Patent  
 RL.P Roles from patents: BIOL (Biological study); USES (Uses)

Absolute stereochemistry.

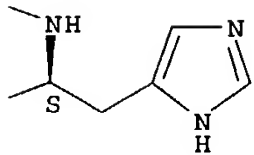
PAGE 1-A



PAGE 1-B



PAGE 1-C

-NH2

\*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

1 REFERENCES IN FILE CA (1907 TO DATE)  
 1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

L3 ANSWER 4 OF 11 REGISTRY COPYRIGHT 2004 ACS on STN  
 RN 373389-66-3 REGISTRY  
 CN Phosphatase, protein phosphotyrosine, SL (9CI) (CA INDEX NAME)

OTHER NAMES:

CN Protein tyrosine phosphatase SL  
 CN Protein tyrosine phosphatase, receptor type Q

CN PTP-SL

CN PTPBR7

CN PTPRQ

MF Unspecified

CI MAN

SR CA

LC STN Files: BIOSIS, CA, CAPLUS, TOXCENTER, USPATFULL

DT.CA CAplus document type: Journal; Patent

RL.P Roles from patents: ANST (Analytical study); BIOL (Biological study);  
 RACT (Reactant or reagent); USES (Uses)RL.NP Roles from non-patents: BIOL (Biological study); PROC (Process); PRP  
 (Properties)

\*\*\* STRUCTURE DIAGRAM IS NOT AVAILABLE \*\*\*

13 REFERENCES IN FILE CA (1907 TO DATE)  
 13 REFERENCES IN FILE CAPLUS (1907 TO DATE)

L3 ANSWER 5 OF 11 REGISTRY COPYRIGHT 2004 ACS on STN

RN 148851-08-5 REGISTRY

CN L-Leucine, L-.alpha.-glutamyl-L-asparaginyl-L-.alpha.-aspartyl-O-phosphono-L-tyrosyl-L-isoleucyl-L-asparaginyl-L-alanyl-L-seryl- (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN L-Leucine, N-[N-[N-[N2-[N-[N-(N2-L-.alpha.-glutamyl-L-asparaginyl)-L-.alpha.-aspartyl]-O-phosphono-L-tyrosyl]-L-isoleucyl]-L-asparaginyl]-L-alanyl]-L-seryl]-

OTHER NAMES:

CN 1: PN: EP1201764 PAGE: 13 claimed protein

FS PROTEIN SEQUENCE; STEREOSEARCH

MF C44 H68 N11 O21 P

SR CA

LC STN Files: CA, CAPLUS, CHEMCATS, TOXCENTER, USPATFULL

DT.CA CAplus document type: Journal; Patent

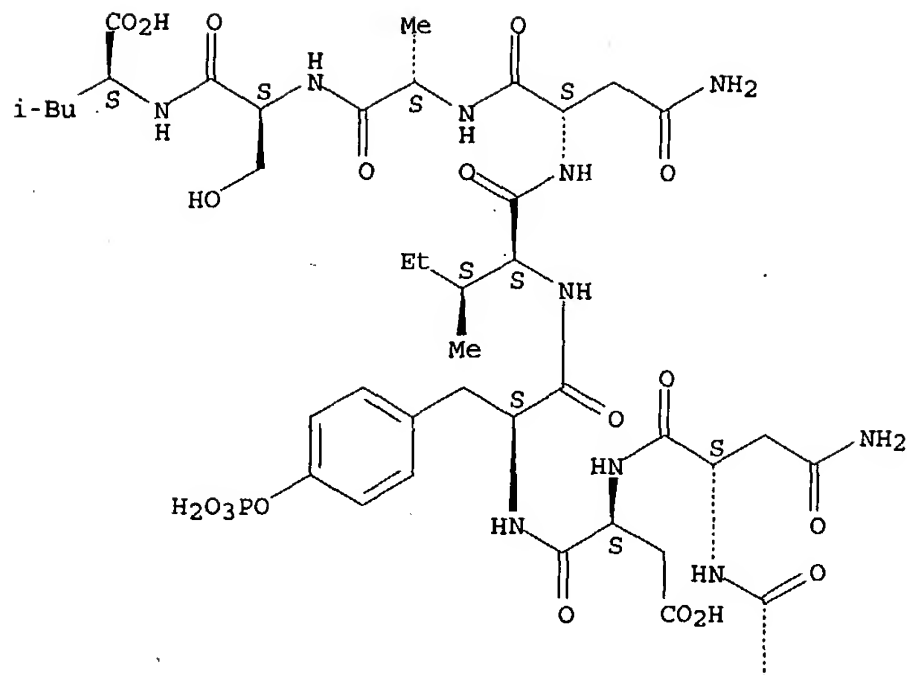
RL.P Roles from patents: ANST (Analytical study); BIOL (Biological study);  
 PROC (Process); USES (Uses)

RL.NP Roles from non-patents: ANST (Analytical study); BIOL (Biological study); FORM (Formation, nonpreparative); PREP (Preparation)

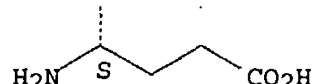
\*\*RELATED SEQUENCES AVAILABLE WITH SEQLINK\*\*

Absolute stereochemistry.

PAGE 1-A



PAGE 2-A



5 REFERENCES IN FILE CA (1907 TO DATE)

5 REFERENCES IN FILE CAPLUS (1907 TO DATE)

L3 ANSWER 6 OF 11 REGISTRY COPYRIGHT 2004 ACS on STN  
 RN 142243-02-5 REGISTRY  
 CN Kinase (phosphorylating), mitogen-activated protein (9CI) (CA INDEX NAME)  
 OTHER NAMES:  
 CN ERK  
 CN ERK kinase  
 CN Erk receptor tyrosine kinase  
 CN ERK/MAP kinase  
 CN Extracellular signal-regulated kinase  
 CN Extracellular signal-regulated protein kinase  
 CN Gene ERK protein kinase  
 CN MAP kinase  
 CN MAP/ERK kinase  
 CN MAPK  
 CN Mitogen-activated protein kinase  
 CN p43 MAP kinase  
 CN p43 Mitogen-activated protein kinase  
 CN p45 MAP kinase  
 DR 133876-94-5, 141349-99-7, 141350-00-7, 141616-09-3  
 MF Unspecified  
 CI MAN  
 SR CA  
 LC STN Files: ADISNEWS, AGRICOLA, BIOPHARMA, BIOSIS, BIOTECHNO, CA, CAPLUS, CEN, CIN, EMBASE, PROMT, TOXCENTER, USPAT2, USPATFULL  
 DT.CA CAplus document type: Book; Conference; Dissertation; Journal; Patent; Report  
 RL.P Roles from patents: ANST (Analytical study); BIOL (Biological study); FORM (Formation, nonpreparative); MSC (Miscellaneous); OCCU (Occurrence); PREP (Preparation); PROC (Process); PRP (Properties); RACT (Reactant or reagent); USES (Uses)  
 RLD.P Roles for non-specific derivatives from patents: ANST (Analytical study); BIOL (Biological study); PREP (Preparation); PROC (Process); PRP (Properties); USES (Uses)  
 RL.NP Roles from non-patents: ANST (Analytical study); BIOL (Biological study); FORM (Formation, nonpreparative); MSC (Miscellaneous); OCCU (Occurrence); PREP (Preparation); PROC (Process); PRP (Properties); RACT (Reactant or reagent); USES (Uses)

RLD.NP Roles for non-specific derivatives from non-patents: ANST (Analytical study); BIOL (Biological study); OCCU (Occurrence); PREP (Preparation); PROC (Process); PRP (Properties); USES (Uses)

\*\*\* STRUCTURE DIAGRAM IS NOT AVAILABLE \*\*\*

9018 REFERENCES IN FILE CA (1907 TO DATE)

55 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA

9063 REFERENCES IN FILE CAPLUS (1907 TO DATE)

L3 ANSWER 7 OF 11 REGISTRY COPYRIGHT 2004 ACS on STN

RN 50812-37-8 REGISTRY

CN Transferase, glutathione S- (9CI) (CA INDEX NAME)

OTHER NAMES:

CN 173: PN: US20040058881 PAGE: 31 claimed sequence

CN 1: PN: US20040146861 FIGURE: 2 claimed sequence

CN 80: PN: WO2004025259 PAGE: 54 claimed sequence

CN 88: PN: WO2004025259 PAGE: 57 claimed sequence

CN Alkyltransferase, glutathione S-

CN Aralkyltransferase, glutathione S-

CN Aryltransferase, glutathione S-

CN Bromosulfophthalein glutathione transferase

CN E.C. 1.8.6.1

CN E.C. 2.5.1.12

CN E.C. 2.5.1.13

CN E.C. 2.5.1.14

CN E.C. 2.5.1.18

CN E.C. 4.4.1.7

CN Epoxidetransferase, glutathione S-

CN Fosfomycin:glutathione S-transferase

CN Glutathione S-alkyltransferase

CN Glutathione S-aralkyltransferase

CN Glutathione S-aralkyltransferase

CN Glutathione S-aryltransferase

CN Glutathione S-epoxidetransferase

CN Glutathione S-methyltransferase

CN Glutathione S-transferase

CN Glutathione S-transferase

CN Glutathione S-transferase .zeta.

CN Glutathione transferase

CN glutathione-S-transferase

CN Glutathionyl transferase

CN GSH S-aryltransferase

CN GSH transferase

CN Ligandins

CN Lyase, hydroxyalkylglutathione

CN Reductase, nitrate ester

CN S-(Hydroxyalkyl)glutathione lyase

CN Thiadiazolidine isomerase

DR 9029-41-8, 9052-42-0, 9079-09-8, 51570-22-0, 37277-81-9, 37290-93-0

MF Unspecified

CI MAN

LC STN Files: ADISNEWS, AGRICOLA, ANABSTR, BIOBUSINESS, BIOSIS, BIOTECHNO, CA, CABA, CAPLUS, CEN, CHEMCATS, CIN, CSCHEM, CSNB, EMBASE, MSDS-OHS, NAPRALERT, NIOSHTIC, PIRA, PROMT, TOXCENTER, ULIDAT, USPAT2, USPATFULL

DT.CA CAplus document type: Book; Conference; Dissertation; Journal; Patent; Preprint; Report

RL.P Roles from patents: ANST (Analytical study); BIOL (Biological study); FORM (Formation, nonpreparative); MSC (Miscellaneous); OCCU (Occurrence); PREP (Preparation); PROC (Process); PRP (Properties); RACT (Reactant or reagent); USES (Uses)

RLD.P Roles for non-specific derivatives from patents: ANST (Analytical study); BIOL (Biological study); CMBI (Combinatorial study); FORM (Formation, nonpreparative); OCCU (Occurrence); PREP (Preparation); PROC (Process); PRP (Properties); RACT (Reactant or reagent); USES (Uses)

RL.NP Roles from non-patents: ANST (Analytical study); BIOL (Biological study); CMBI (Combinatorial study); FORM (Formation, nonpreparative); MSC (Miscellaneous); OCCU (Occurrence); PREP (Preparation); PROC (Process); PRP (Properties); RACT (Reactant or reagent); USES (Uses); NORL (No role in record)

RLD.NP Roles for non-specific derivatives from non-patents: ANST (Analytical study); BIOL (Biological study); FORM (Formation, nonpreparative); OCCU (Occurrence); PREP (Preparation); PROC (Process); PRP (Properties); RACT (Reactant or reagent); USES (Uses)

\*\*\* STRUCTURE DIAGRAM IS NOT AVAILABLE \*\*\*

14626 REFERENCES IN FILE CA (1907 TO DATE)

781 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA  
14678 REFERENCES IN FILE CAPLUS (1907 TO DATE)

L3 ANSWER 8 OF 11 REGISTRY COPYRIGHT 2004 ACS on STN  
 RN 13721-39-6 REGISTRY  
 CN Sodium vanadium oxide (Na<sub>3</sub>VO<sub>4</sub>) (9CI) (CA INDEX NAME)  
 OTHER CA INDEX NAMES:  
 CN Sodium vanadate(V) (Na<sub>3</sub>VO<sub>4</sub>) (6CI)  
 CN Vanadic acid (H<sub>3</sub>VO<sub>4</sub>), trisodium salt (8CI)  
 OTHER NAMES:  
 CN NSC 79534  
 CN Sodium o-vanadate  
 CN Sodium orthovanadate  
 CN Sodium orthovanadate (Na<sub>3</sub>VO<sub>4</sub>)  
 CN Sodium tetraoxovanadate(3-)  
 CN Sodium vanadate  
 CN Sodium vanadate (Na<sub>3</sub>VO<sub>4</sub>)  
 CN Trisodium orthovanadate  
 CN Trisodium vanadate  
 CN Vanadate (VO<sub>4</sub>3-), trisodium, (T-4)-  
 DR 15066-70-3, 70904-59-5  
 MF Na . O . V  
 AF Na<sub>3</sub> O<sub>4</sub> V  
 CI COM, TIS  
 LC STN Files: AGRICOLA, AQUIRE, BIOBUSINESS, BIOSIS, BIOTECHNO, CA, CAOLD, CAPLUS, CASREACT, CHEMCATS, CHEMLIST, CIN, CSCHEM, CSNB, DDFU, DRUGU, EMBASE, IFICDB, IFIPAT, IFIUDB, MSDS-OHS, NIOSHTIC, PIRA, PROMT, RTECS\*, TOXCENTER, TULSA, ULIDAT, USPAT2, USPATFULL  
 (\*File contains numerically searchable property data)  
 Other Sources: DSL\*\*, EINECS\*\*, TSCA\*\*  
 (\*\*Enter CHEMLIST File for up-to-date regulatory information)  
 DT.CA Cplus document type: Conference; Dissertation; Journal; Patent; Report  
 RL.P Roles from patents: ANST (Analytical study); BIOL (Biological study); OCCU (Occurrence); PREP (Preparation); PROC (Process); PRP (Properties); RACT (Reactant or reagent); USES (Uses); NORL (No role in record)  
 RLD.P Roles for non-specific derivatives from patents: BIOL (Biological study); PREP (Preparation); USES (Uses)  
 RL.NP Roles from non-patents: ANST (Analytical study); BIOL (Biological study); FORM (Formation, nonpreparative); OCCU (Occurrence); PREP (Preparation); PROC (Process); PRP (Properties); RACT (Reactant or reagent); USES (Uses); NORL (No role in record)  
 RLD.NP Roles for non-specific derivatives from non-patents: BIOL (Biological study); PRP (Properties)

Component	Ratio	Component Registry Number
O	4	17778-80-2
V	1	7440-62-2
Na	3	7440-23-5

879 REFERENCES IN FILE CA (1907 TO DATE)  
 8 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA  
 883 REFERENCES IN FILE CAPLUS (1907 TO DATE)  
 21 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

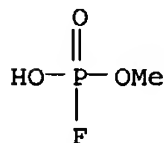
L3 ANSWER 9 OF 11 REGISTRY COPYRIGHT 2004 ACS on STN  
 RN 9061-61-4 REGISTRY  
 CN Nerve growth factor (9CI) (CA INDEX NAME)  
 OTHER NAMES:  
 CN Nerve growth hormone  
 CN NGF  
 MF Unspecified  
 CI PMS, COM, MAN  
 PCT Manual registration  
 LC STN Files: ADISINSIGHT, AGRICOLA, ANABSTR, BIOBUSINESS, BIOSIS, BIOTECHNO, CA, CANCERLIT, CAPLUS, CBNB, CEN, CHEMCATS, CIN, CSCHEM, DDFU, DRUGU, EMBASE, IFICDB, IFIPAT, IFIUDB, IPA, MEDLINE, MRCK\*, PHAR, PROMT, TOXCENTER, USPAT2, USPATFULL  
 (\*File contains numerically searchable property data)  
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 RL.P Roles from patents: ANST (Analytical study); BIOL (Biological study); FORM (Formation, nonpreparative); MSC (Miscellaneous); OCCU (Occurrence); PREP (Preparation); PROC (Process); PRP (Properties); RACT (Reactant or reagent); USES (Uses); NORL (No role in record)

RLD.P Roles for non-specific derivatives from patents: ANST (Analytical study); BIOL (Biological study); PREP (Preparation); PROC (Process); PRP (Properties); RACT (Reactant or reagent); USES (Uses)  
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\*\*\* STRUCTURE DIAGRAM IS NOT AVAILABLE \*\*\*

10172 REFERENCES IN FILE CA (1907 TO DATE)  
 137 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA  
 10195 REFERENCES IN FILE CAPLUS (1907 TO DATE)

L3 ANSWER 10 OF 11 REGISTRY COPYRIGHT 2004 ACS on STN  
 RN 754-02-9 REGISTRY  
 CN Phosphorofluoridic acid, monomethyl ester (9CI) (CA INDEX NAME)  
 OTHER CA INDEX NAMES:  
 CN Methyl phosphorofluoridate ((MeO)(HO)FPO) (6CI, 7CI)  
 FS 3D CONCORD  
 MF C H4 F O3 P  
 CI COM  
 LC STN Files: BEILSTEIN\*, CA, CAOLD, CAPLUS, TOXCENTER, USPATFULL  
 (\*File contains numerically searchable property data)  
 DT.CA CAplus document type: Journal; Patent  
 RL.P Roles from patents: BIOL (Biological study); USES (Uses)  
 RL.NP Roles from non-patents: NORL (No role in record)



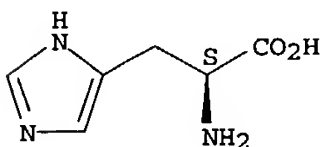
\*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

4 REFERENCES IN FILE CA (1907 TO DATE)  
 4 REFERENCES IN FILE CAPLUS (1907 TO DATE)  
 3 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

L3 ANSWER 11 OF 11 REGISTRY COPYRIGHT 2004 ACS on STN  
 RN 71-00-1 REGISTRY  
 CN L-Histidine (9CI) (CA INDEX NAME)  
 OTHER CA INDEX NAMES:  
 CN Histidine, L- (8CI)  
 OTHER NAMES:  
 CN (S)-.alpha.-Amino-1H-imidazole-4-propanoic acid  
 CN (S)-4-(2-Amino-2-carboxyethyl)imidazole  
 CN (S)-Histidine  
 CN 1H-Imidazole-4-alanine, (S)-  
 CN 1H-Imidazole-4-propanoic acid, .alpha.-amino-, (S)-  
 CN Glyoxaline-5-alanine  
 CN Histidine  
 CN L-(-)-Histidine  
 CN L-Alanine, 3-(1H-imidazol-4-yl)-  
 CN NSC 137773  
 FS STEREOSEARCH  
 DR 7006-35-1, 150-35-6, 54166-13-1, 155304-24-8, 35479-49-3, 35558-59-9,  
 45955-20-2  
 MF C6 H9 N3 O2  
 CI COM  
 LC STN Files: ADISNEWS, AGRICOLA, ANABSTR, BEILSTEIN\*, BIOBUSINESS, BIOSIS,  
 BIOTECHNO, CA, CABA, CANCERLIT, CAOLD, CAPLUS, CASREACT, CBNB, CEN,  
 CHEMCATS, CHEMINFORMRX, CHEMLIST, CIN, CSCHEM, CSNB, DDFU, DETHERM\*,  
 DIOGENES, DRUGU, EMBASE, GMELIN\*, HODOC\*, HSDB\*, IFICDB, IFIPAT, IFIUDB,  
 IPA, MEDLINE, MRCK\*, MSDS-OHS, NAPRALERT, NIOSHTIC, PIRA, PROMT, PS,  
 RTECS\*, SPECINFO, TOXCENTER, TULSA, ULIDAT, USAN, USPAT2, USPATFULL  
 (\*File contains numerically searchable property data)  
 Other Sources: DSL\*\*, EINECS\*\*, TSCA\*\*, WHO  
 (\*\*Enter CHEMLIST File for up-to-date regulatory information)

DT.CA CAplus document type: Book; Conference; Dissertation; Journal; Patent; Preprint; Report  
 RL.P Roles from patents: ANST (Analytical study); BIOL (Biological study); CMBI (Combinatorial study); FORM (Formation, nonpreparative); MSC (Miscellaneous); OCCU (Occurrence); PREP (Preparation); PROC (Process); PRP (Properties); RACT (Reactant or reagent); USES (Uses); NORL (No role in record)  
 RLD.P Roles for non-specific derivatives from patents: ANST (Analytical study); BIOL (Biological study); OCCU (Occurrence); PREP (Preparation); PROC (Process); PRP (Properties); RACT (Reactant or reagent); USES (Uses)  
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 RLD.NP Roles for non-specific derivatives from non-patents: ANST (Analytical study); BIOL (Biological study); FORM (Formation, nonpreparative); MSC (Miscellaneous); OCCU (Occurrence); PREP (Preparation); PROC (Process); PRP (Properties); RACT (Reactant or reagent); USES (Uses)

Absolute stereochemistry. Rotation (-).



\*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

34128 REFERENCES IN FILE CA (1907 TO DATE)  
 1379 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA  
 34185 REFERENCES IN FILE CAPLUS (1907 TO DATE)  
 5 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

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FILE 'WPIX' ENTERED AT 09:59:08 ON 30 SEP 2004  
 COPYRIGHT (C) 2004 THE THOMSON CORPORATION

FILE LAST UPDATED: 28 SEP 2004 <20040928/UP>  
 MOST RECENT DERWENT UPDATE: 200462 <200462/DW>  
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>>> FOR DETAILS OF THE PATENTS COVERED IN CURRENT UPDATES, SEE  
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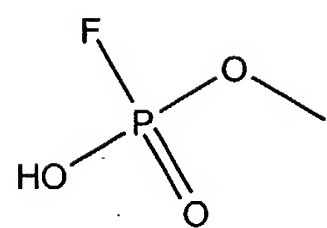
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L4 ANSWER 1 OF 1 WPIX COPYRIGHT 2004 THE THOMSON CORP on STN  
 AN 2002-418778 [45] WPIX  
 DNC C2002-118290  
 TI Identifying an agent that inhibits the catalytic activity of a tyrosine phosphatase, for treating neurodegenerative diseases, comprises quantitating dephosphorylation of a substrate by the enzyme, in the presence of the test agent.

DC B04 D16  
 IN SCHAEFFER, E  
 PA (PFIZ) PFIZER PROD INC; (SCHA-I) SCHAEFFER E  
 CYC 29  
 PI EP 1201764 A2 20020502 (200245)\* EN 14 C12Q001-42  
 R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT  
 RO SE SI TR  
 US 2002086342 A1 20020704 (200247) C12Q001-48 <--  
 CA 2360049 A1 20020430 (200248) EN C12Q001-42  
 JP 2002355066 A 20021210 (200311) 15 C12N015-09  
 ADT EP 1201764 A2 EP 2001-124850 20011018; US 2002086342 A1 Provisional US  
 2000-244539P 20001031, US 2001-437 20011031; CA 2360049 A1 CA 2001-2360049  
 20011029; JP 2002355066 A JP 2001-334245 20011031  
 PRAI US 2000-244539P 20001031; US 2001-437 20011031  
 IC ICM C12N015-09; C12Q001-42; C12Q001-48  
 ICS A61K038-43; A61K045-00; A61P025-00; A61P025-28; A61P043-00;  
 C12Q001-00; C12Q001-02; C12Q001-44; G01N021-78; G01N033-15;  
 G01N033-50; G01N033-573  
 AB EP 1201764 A UPAB: 20020717  
 NOVELTY - Identifying an agent that inhibits the catalytic activity of protein tyrosine phosphatase, PTPbr7, comprising quantitating and comparing the dephosphorylation in two cocktails, one containing PTPbr7, a substrate that can be dephosphorylated by PTPbr7 and a reducing buffer, and the other having the same ingredients but lacking the test agent.  
 DETAILED DESCRIPTION - Identifying an agent that inhibits the catalytic activity of protein tyrosine phosphatase, PTPbr7, comprising:  
 (a) combining, in a first cocktail, PTPbr7, a substrate capable of being dephosphorylated by PTPbr7, and a test agent, in an assay buffer containing a reducing buffer;  
 (b) preparing a second cocktail comprising all the ingredients of the first cocktail except for the test agent;  
 (c) incubating the first and second cocktails to allow dephosphorylation of the substrate by PTPbr7;  
 (d) quantitating the dephosphorylation in each of the cocktails; and  
 (e) comparing the amounts of dephosphorylation, where a PTPbr7 inhibitor is a test agent whose presence results in less dephosphorylation than its absence.  
 INDEPENDENT CLAIMS are also included for the following:  
 (1) a pharmaceutical composition comprising an inhibitor of the catalytic activity of PTPbr7; and  
 (2) screening for an agent that inhibits the catalytic activity of PTPbr7, comprising:  
 (a) exposing cells or a cell line expressing PTPbr7 and capable of responding to nerve growth factor (NGF), to NGF, in the presence and absence of a test agent, to allow for a NGF response to occur in the presence of a PTPbr7 inhibitor;  
 (b) detecting the response; and  
 (c) comparing the response, where the PTPbr7 inhibitor is a test agent whose presence results in more of an NGF response than in its absence.  
 ACTIVITY - Nootropic; Neuroprotective. No suitable biological data is given.  
 MECHANISM OF ACTION - PTPbr7 inhibitor.  
 USE - The method is used to identify an agent that inhibits the catalytic activity of PTPbr7 (claimed). The agent can be used to treat neurodegenerative diseases.  
 Dwg.0/0  
 FS CPI  
 FA AB; DCN  
 MC CPI: B04-F01; B04-M01; B04-N08; B11-C07B2; B12-K04E; B14-J01A; D05-H09

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=>



o-methylfluorophosphate



=> b reg  
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USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT.  
PLEASE SEE "HELP USAGETERMS" FOR DETAILS.  
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STRUCTURE FILE UPDATES: 29 SEP 2004 HIGHEST RN 754169-63-6  
DICTIONARY FILE UPDATES: 29 SEP 2004 HIGHEST RN 754169-63-6

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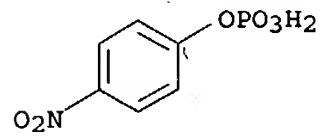
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conducting SmartSELECT searches.

Crossover limits have been increased. See HELP CROSSOVER for details.

Experimental and calculated property data are now available. For more  
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<http://www.cas.org/ONLINE/DBSS/registryss.html>

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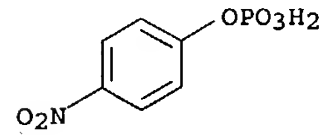
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RN 333338-18-4 REGISTRY  
CN Phosphoric acid, mono(4-nitrophenyl) ester, disodium salt,  
hexahydrate (9CI) (CA INDEX NAME)  
MF C6 H6 N O6 P . 6 H2 O . 2 Na  
SR CAS Client Services  
LC STN Files: CHEMCATS  
CRN (330-13-2)



● 2 Na

● 6 H<sub>2</sub>O

L31 ANSWER 2 OF 24 REGISTRY COPYRIGHT 2004 ACS on STN  
RN 208651-58-5 REGISTRY  
CN Phosphoric acid, mono(4-nitrophenyl) ester, monopotassium salt  
(9CI) (CA INDEX NAME)  
MF C6 H6 N O6 P . K  
SR CAS Client Services  
LC STN Files: CA, CAPLUS, CASREACT, CHEMCATS, CSCHEM  
DT.CA CAplus document type: Journal  
RL.NP Roles from non-patents: PREP (Preparation)  
CRN (330-13-2)

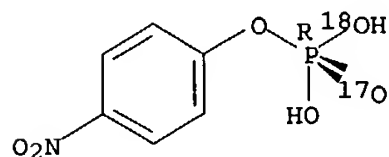


● K

1 REFERENCES IN FILE CA (1907 TO DATE)  
 1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

L31 ANSWER 3 OF 24 REGISTRY COPYRIGHT 2004 ACS on STN  
 RN 112115-01-2 REGISTRY  
 CN Phosphoric-170-180 acid, 160-(4-nitrophenyl) ester, disodium salt,  
 (R)- (9CI) (CA INDEX NAME)  
 FS STEREOSEARCH  
 MF C6 H6 N O6 P . 2 Na  
 SR CA  
 LC STN Files: CA, CAPLUS  
 DT.CA CAplus document type: Journal  
 RL.NP Roles from non-patents: PREP (Preparation)  
 CRN (112114-76-8)

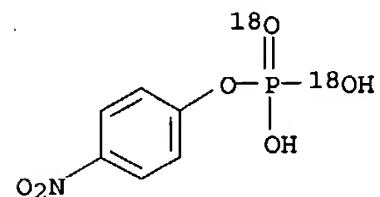
Absolute stereochemistry.



●2 Na

1 REFERENCES IN FILE CA (1907 TO DATE)  
 1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

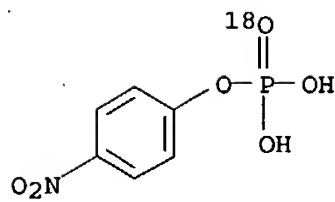
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 (9CI) (CA INDEX NAME)  
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 SR CA  
 LC STN Files: CA, CAPLUS  
 DT.CA CAplus document type: Journal  
 RL.NP Roles from non-patents: PREP (Preparation)



●2 Na

1 REFERENCES IN FILE CA (1907 TO DATE)  
 1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

L31 ANSWER 5 OF 24 REGISTRY COPYRIGHT 2004 ACS on STN  
 RN 112114-99-5 REGISTRY  
 CN Phosphoric-180 acid, 160-(4-nitrophenyl) ester, disodium salt  
 (9CI) (CA INDEX NAME)  
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 DT.CA CAplus document type: Journal  
 RL.NP Roles from non-patents: PREP (Preparation)

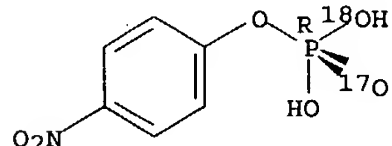


● 2 Na

1 REFERENCES IN FILE CA (1907 TO DATE)  
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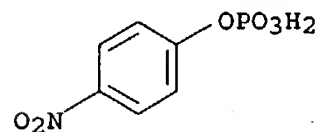
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 RN 112114-76-8 REGISTRY  
 CN Phosphoric-170-180 acid, 160-(4-nitrophenyl) ester, (R)- (9CI)  
 (CA INDEX NAME)  
 FS STEREOSEARCH  
 MF C6 H6 N O6 P  
 CI COM  
 SR CA  
 LC STN Files: CA, CAPLUS  
 DT.CA CAplus document type: Journal  
 RL.NP Roles from non-patents: BIOL (Biological study); PREP (Preparation)

Absolute stereochemistry.



2 REFERENCES IN FILE CA (1907 TO DATE)  
 2 REFERENCES IN FILE CAPLUS (1907 TO DATE)

L31 ANSWER 7 OF 24 REGISTRY COPYRIGHT 2004 ACS on STN  
 RN 110347-83-6 REGISTRY  
 CN Phosphoric acid, mono(4-nitrophenyl) ester, magnesium salt (1:1)  
 (9CI) (CA INDEX NAME)  
 MF C6 H6 N O6 P . Mg  
 SR CA  
 LC STN Files: CA, CAPLUS  
 DT.CA CAplus document type: Journal  
 RL.NP Roles from non-patents: BIOL (Biological study); RACT (Reactant or reagent)  
 CRN (330-13-2)

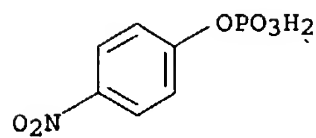


● Mg

2 REFERENCES IN FILE CA (1907 TO DATE)  
 2 REFERENCES IN FILE CAPLUS (1907 TO DATE)

L31 ANSWER 8 OF 24 REGISTRY COPYRIGHT 2004 ACS on STN  
 RN 88948-42-9 REGISTRY  
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 (CA INDEX NAME)  
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 DT.CA CAplus document type: Journal  
 RL.NP Roles from non-patents: RACT (Reactant or reagent)

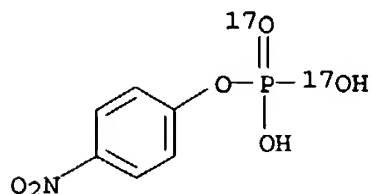
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●x Ca

1 REFERENCES IN FILE CA (1907 TO DATE)  
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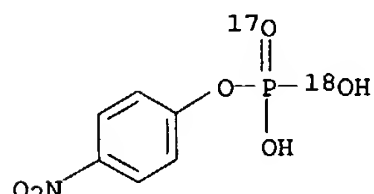
L31 ANSWER 9 OF 24 REGISTRY COPYRIGHT 2004 ACS on STN  
 RN 87174-81-0 REGISTRY  
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 LC STN Files: CA, CAPLUS  
 DT.CA CAplus document type: Journal  
 RL.NP Roles from non-patents: PRP (Properties)



●2 Na

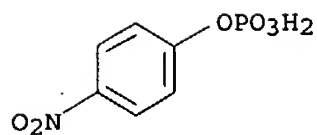
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 1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

L31 ANSWER 10 OF 24 REGISTRY COPYRIGHT 2004 ACS on STN  
 RN 81939-55-1 REGISTRY  
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 MF C6 H6 N O6 P  
 LC STN Files: CA, CAPLUS  
 DT.CA CAplus document type: Journal  
 RL.NP Roles from non-patents: FORM (Formation, nonpreparative)



1 REFERENCES IN FILE CA (1907 TO DATE)  
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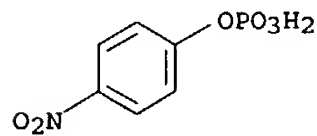
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 RN 77849-47-9 REGISTRY  
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 MF C6 H6 N O6 P . 1/2 Zr  
 LC STN Files: CA, CAPLUS, USPATFULL  
 DT.CA CAplus document type: Patent  
 RL.P Roles from patents: PREP (Preparation)  
 CRN (330-13-2)



●1/2 Zr(IV)

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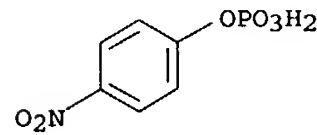
L31 ANSWER 12 OF 24 REGISTRY COPYRIGHT 2004 ACS on STN  
 RN 76019-14-2 REGISTRY  
 CN Phosphoric acid, mono(4-nitrophenyl) ester, magnesium salt (9CI)  
 (CA INDEX NAME)  
 MF C<sub>6</sub> H<sub>6</sub> N O<sub>6</sub> P . x Mg  
 LC STN Files: CA, CAPLUS  
 DT.CA Caplus document type: Journal; Patent  
 RL.P Roles from patents: BIOL (Biological study)  
 RL.NP Roles from non-patents: RACT (Reactant or reagent)  
 CRN (330-13-2)



●x Mg

3 REFERENCES IN FILE CA (1907 TO DATE)  
 3 REFERENCES IN FILE CAPLUS (1907 TO DATE)

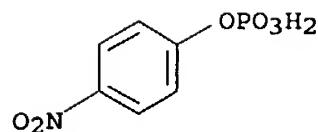
L31 ANSWER 13 OF 24 REGISTRY COPYRIGHT 2004 ACS on STN  
 RN 75431-32-2 REGISTRY  
 CN Phosphoric acid, mono(4-nitrophenyl) ester, thorium(4+) salt (2:1)  
 (9CI) (CA INDEX NAME)  
 MF C<sub>6</sub> H<sub>6</sub> N O<sub>6</sub> P . 1/2 Th  
 LC STN Files: CA, CAPLUS, USPATFULL  
 DT.CA Caplus document type: Journal; Patent  
 RL.P Roles from patents: PREP (Preparation)  
 RL.NP Roles from non-patents: PREP (Preparation)  
 CRN (330-13-2)



●1/2 Th(IV)

3 REFERENCES IN FILE CA (1907 TO DATE)  
 3 REFERENCES IN FILE CAPLUS (1907 TO DATE)

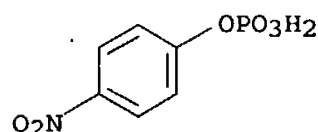
L31 ANSWER 14 OF 24 REGISTRY COPYRIGHT 2004 ACS on STN  
 RN 71735-29-0 REGISTRY  
 CN Phosphoric acid, mono(4-nitrophenyl) ester, disodium salt, hydrate  
 (9CI) (CA INDEX NAME)  
 MF C<sub>6</sub> H<sub>6</sub> N O<sub>6</sub> P . x H<sub>2</sub>O . 2 Na  
 LC STN Files: BEILSTEIN\*  
 (\*File contains numerically searchable property data)  
 CRN (330-13-2)



● 2 Na

● x H<sub>2</sub>O

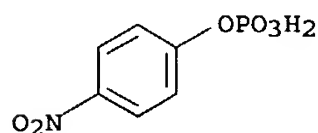
L31 ANSWER 15 OF 24 REGISTRY COPYRIGHT 2004 ACS on STN  
 RN 70951-20-1 REGISTRY  
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 (9CI) (CA INDEX NAME)  
 MF C<sub>6</sub> H<sub>6</sub> N O<sub>6</sub> P . Ca  
 LC STN Files: CA, CAPLUS  
 DT.CA CAplus document type: Journal  
 RL.NP Roles from non-patents: RACT (Reactant or reagent)  
 CRN (330-13-2)



● Ca

1 REFERENCES IN FILE CA (1907 TO DATE)  
 1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

L31 ANSWER 16 OF 24 REGISTRY COPYRIGHT 2004 ACS on STN  
 RN 63035-79-0 REGISTRY  
 CN Phosphoric acid, mono(4-nitrophenyl) ester, barium salt (1:1)  
 (9CI) (CA INDEX NAME)  
 MF C<sub>6</sub> H<sub>6</sub> N O<sub>6</sub> P . Ba  
 LC STN Files: CA, CAPLUS  
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 CRN (330-13-2)

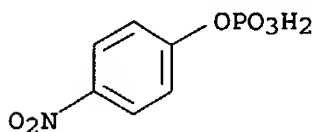


● Ba

12 REFERENCES IN FILE CA (1907 TO DATE)  
 12 REFERENCES IN FILE CAPLUS (1907 TO DATE)

L31 ANSWER 17 OF 24 REGISTRY COPYRIGHT 2004 ACS on STN  
 RN 54306-27-3 REGISTRY  
 CN Phosphoric acid, mono(4-nitrophenyl) ester, sodium salt (9CI)  
 (CA INDEX NAME)  
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 LC STN Files: BEILSTEIN\*, CA, CAPLUS, CHEMLIST  
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 Other Sources: EINECS\*\*, NDSL\*\*, TSCA\*\*  
 (\*\*Enter CHEMLIST File for up-to-date regulatory information)

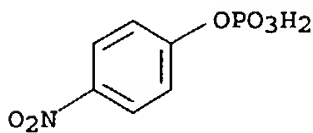
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 RL.P Roles from patents: PREP (Preparation)  
 CRN (330-13-2)



● x Na

1 REFERENCES IN FILE CA (1907 TO DATE)  
 1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

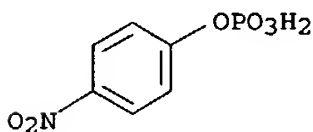
L31 ANSWER 18 OF 24 REGISTRY COPYRIGHT 2004 ACS on STN  
 RN 51952-77-3 REGISTRY  
 CN Phosphoric acid, mono(4-nitrophenyl) ester, disilver(1+) salt (9CI) (CA INDEX NAME)  
 MF C6 H6 N O6 P . 2 Ag  
 LC STN Files: BEILSTEIN\*, CA, CAPLUS  
 (\*File contains numerically searchable property data)  
 DT.CA CAplus document type: Journal; Patent  
 RL.P Roles from patents: RACT (Reactant or reagent)  
 RL.NP Roles from non-patents: PREP (Preparation); RACT (Reactant or reagent)  
 CRN (330-13-2)



● 2 Ag(I)

2 REFERENCES IN FILE CA (1907 TO DATE)  
 2 REFERENCES IN FILE CAPLUS (1907 TO DATE)

L31 ANSWER 19 OF 24 REGISTRY COPYRIGHT 2004 ACS on STN  
 RN 32348-91-7 REGISTRY  
 CN Phosphoric acid, mono(4-nitrophenyl) ester, diammonium salt (9CI) (CA INDEX NAME)  
 OTHER NAMES:  
 CN Diammonium 4-nitrophenyl phosphate  
 CN Diammonium p-nitrophenyl phosphate  
 MF C6 H6 N O6 P . 2 H3 N  
 LC STN Files: BEILSTEIN\*, CA, CAPLUS, CHEMLIST, IFICDB, IFIPAT, IFIUDB, USPATFULL  
 (\*File contains numerically searchable property data)  
 Other Sources: EINECS\*\*, NDSL\*\*, TSCA\*\*  
 (\*\*Enter CHEMLIST File for up-to-date regulatory information)  
 DT.CA CAplus document type: Patent  
 RL.P Roles from patents: BIOL (Biological study); PROC (Process)  
 CRN (330-13-2)

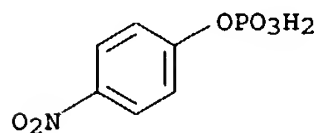


● 2 NH3

3 REFERENCES IN FILE CA (1907 TO DATE)

## 3 REFERENCES IN FILE CAPLUS (1907 TO DATE)

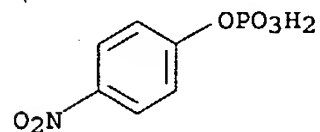
L31 ANSWER 20 OF 24 REGISTRY COPYRIGHT 2004 ACS on STN  
 RN 32348-90-6 REGISTRY  
 CN Phosphoric acid, mono(4-nitrophenyl) ester, magnesium salt (1:2)  
 (9CI) (CA INDEX NAME)  
 OTHER NAMES:  
 CN Dimagnesium p-nitrophenyl phosphate  
 MF C6 H6 N O6 P . 2 Mg  
 LC STN Files: CA, CAPLUS, CHEMCATS, CHEMLIST, IFICDB, IFIPAT, IFIUDB,  
 USPATFULL  
 Other Sources: EINECS\*\*, NDSL\*\*, TSCA\*\*  
 (\*\*Enter CHEMLIST File for up-to-date regulatory information)  
 DT.CA CAplus document type: Patent  
 RL.P Roles from patents: BIOL (Biological study); PROC (Process)  
 CRN (330-13-2)



●2 Mg

2 REFERENCES IN FILE CA (1907 TO DATE)  
 2 REFERENCES IN FILE CAPLUS (1907 TO DATE)

L31 ANSWER 21 OF 24 REGISTRY COPYRIGHT 2004 ACS on STN  
 RN 16785-19-6 REGISTRY  
 CN Phosphoric acid, mono(4-nitrophenyl) ester, dipotassium salt (9CI)  
 (CA INDEX NAME)  
 OTHER CA INDEX NAMES:  
 CN Phosphoric acid, mono(p-nitrophenyl) ester, dipotassium salt (8CI)  
 MF C6 H6 N O6 P . 2 K  
 CRN (330-13-2)



●2 K

L31 ANSWER 22 OF 24 REGISTRY COPYRIGHT 2004 ACS on STN  
 RN 4264-83-9 REGISTRY  
 CN Phosphoric acid, mono(4-nitrophenyl) ester, disodium salt (9CI)  
 (CA INDEX NAME)  
 OTHER CA INDEX NAMES:  
 CN Phenol, p-nitro-, di-H phosphate, disodium salt (7CI)  
 CN Phenol, p-nitro-, phosphate disodium salt (6CI)  
 CN Phosphoric acid, mono(p-nitrophenyl) ester, disodium salt (8CI)  
 OTHER NAMES:  
 CN 4-Nitrophenyl phosphate disodium salt  
 CN Disodium 4-nitrophenyl phosphate  
 CN Disodium mono(4-nitrophenyl) phosphate  
 CN Disodium p-nitrophenyl phosphate  
 CN p-Nitrophenyl disodium phosphate  
 CN p-Nitrophenyl phosphate disodium salt  
 CN p-Nitrophenylphosphate sodium salt  
 CN p-NPP disodium salt  
 MF C6 H6 N O6 P . 2 Na  
 CI COM  
 LC STN Files: AGRICOLA, BEILSTEIN\*, BIOBUSINESS, BIOSIS, CA, CAOLD, CAPLUS,  
 CASREACT, CHEMCATS, CHEMLIST, CSCHEM, IFICDB, IFIPAT, IFIUDB, MSDS-OHS,  
 TOXCENTER, USPATFULL

(\*File contains numerically searchable property data)

Other Sources: DSL\*\*, EINECS\*\*, TSCA\*\*

(\*\*Enter CHEMLIST File for up-to-date regulatory information)

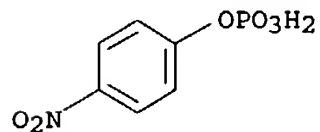
DT.CA CAplus document type: Conference; Journal; Patent

RL.P Roles from patents: ANST (Analytical study); BIOL (Biological study); PREP (Preparation); PROC (Process); RACT (Reactant or reagent); USES (Uses)

RLD.P Roles for non-specific derivatives from patents: ANST (Analytical study)

RL.NP Roles from non-patents: ANST (Analytical study); BIOL (Biological study); OCCU (Occurrence); PREP (Preparation); PROC (Process); PRP (Properties); RACT (Reactant or reagent); USES (Uses); NORL (No role in record)

CRN (330-13-2)



● 2 Na

80 REFERENCES IN FILE CA (1907 TO DATE)

1 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA

80 REFERENCES IN FILE CAPLUS (1907 TO DATE)

6 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

L31 ANSWER 23 OF 24 REGISTRY COPYRIGHT 2004 ACS on STN

RN 4154-43-2 REGISTRY

CN Phosphoric acid, mono(4-nitrophenyl) ester, monosodium salt (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN Phenol, p-nitro-, di-H phosphate, sodium salt (7CI)

CN Phosphoric acid, mono(p-nitrophenyl) ester, monosodium salt (8CI)

OTHER NAMES:

CN Sodium p-nitrophenyl phosphate

MF C6 H6 N O6 P . Na

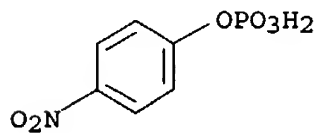
LC STN Files: BEILSTEIN\*, BIOSIS, CA, CAOLD, CAPLUS, CSCHEM, TOXCENTER (\*File contains numerically searchable property data)

DT.CA CAplus document type: Journal; Patent

RL.P Roles from patents: ANST (Analytical study); BIOL (Biological study); USES (Uses)

RL.NP Roles from non-patents: BIOL (Biological study); PREP (Preparation); PROC (Process); PRP (Properties); RACT (Reactant or reagent); NORL (No role in record)

CRN (330-13-2)



● Na

22 REFERENCES IN FILE CA (1907 TO DATE)

22 REFERENCES IN FILE CAPLUS (1907 TO DATE)

2 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

L31 ANSWER 24 OF 24 REGISTRY COPYRIGHT 2004 ACS on STN

RN 330-13-2 REGISTRY

CN Phosphoric acid, mono(4-nitrophenyl) ester (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

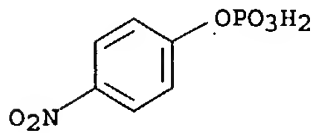
CN Phenol, p-nitro-, dihydrogen phosphate (6CI)

CN Phosphoric acid, mono(p-nitrophenyl) ester (8CI)

CN Phosphoric acid, p-nitrophenyl ester (6CI)

## OTHER NAMES:

CN 4-Nitrophenyl dihydrogen phosphate  
 CN 4-Nitrophenyl phosphate  
 CN NPP  
 CN NSC 404086  
 CN p-Nitrophenol phosphate  
 CN p-Nitrophenyl dihydrogen phosphate  
 CN p-Nitrophenyl phosphate  
 CN PNPP  
 FS 3D CONCORD  
 MF C6 H6 N O6 P  
 CI COM  
 LC STN Files: AGRICOLA; BEILSTEIN\*, BIOBUSINESS, BIOSIS, BIOTECHNO, CA,  
   CANCERLIT, CAOLD, CAPLUS, CASREACT, CHEMCATS, CHEMLIST, CSCHEM, EMBASE,  
   GMELIN\*, IFICDB, IFIPAT, IFIUDB, MEDLINE, MSDS-OHS, NIOSHTIC, PROMT,  
   TOXCENTER, USPATFULL  
   (\*File contains numerically searchable property data)  
 Other Sources: DSL\*\*, EINECS\*\*, TSCA\*\*  
   (\*\*Enter CHEMLIST File for up-to-date regulatory information)  
 DT.CA CAplus document type: Conference; Dissertation; Journal; Patent  
 RL.P Roles from patents: ANST (Analytical study); BIOL (Biological study);  
   PROC (Process); RACT (Reactant or reagent); USES (Uses); NORL (No role  
   in record)  
 RLD.P Roles for non-specific derivatives from patents: ANST (Analytical  
   study); BIOL (Biological study); PREP (Preparation); USES (Uses)  
 RL.NP Roles from non-patents: ANST (Analytical study); BIOL (Biological  
   study); FORM (Formation, nonpreparative); OCCU (Occurrence); PREP  
   (Preparation); PROC (Process); PRP (Properties); RACT (Reactant or  
   reagent); USES (Uses); NORL (No role in record)  
 RLD.NP Roles for non-specific derivatives from non-patents: BIOL (Biological  
   study); FORM (Formation, nonpreparative); PREP (Preparation); PROC  
   (Process); PRP (Properties); RACT (Reactant or reagent)



## \*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

1609 REFERENCES IN FILE CA (1907 TO DATE)  
 24 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA  
 1609 REFERENCES IN FILE CAPLUS (1907 TO DATE)  
 31 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

=> => d his

(FILE 'HOME' ENTERED AT 09:57:54 ON 30 SEP 2004)  
 FILE 'HCAPLUS' ENTERED AT 09:58:04 ON 30 SEP 2004  
 L1       1 US20020086342/PN  
 FILE 'REGISTRY' ENTERED AT 09:58:22 ON 30 SEP 2004  
 FILE 'HCAPLUS' ENTERED AT 09:58:26 ON 30 SEP 2004  
 L2       TRA L1 1- RN :       11 TERMS  
 FILE 'REGISTRY' ENTERED AT 09:58:27 ON 30 SEP 2004  
 L3       11 SEA L2  
 FILE 'WPIX' ENTERED AT 09:58:30 ON 30 SEP 2004  
 L4       1 US20020086342/PN  
 FILE 'REGISTRY' ENTERED AT 10:15:37 ON 30 SEP 2004  
   E PTPBR7/CN  
 L5       1 L3 AND PTPBR7  
 L6       4 PTPBR7  
 FILE 'HCAPLUS' ENTERED AT 10:18:12 ON 30 SEP 2004  
 L7       13 L5  
 L8       5952 (PHOSPHATASE (2A) PROTEIN (2A) (PHOSPHOTYROSINE OR TYROSINE) OR

E DESPHOSPHORYLATION, BIOLOGICAL/CT  
 E DEPHOSPHORYLATION, BIOLOGICAL/CT  
 E DEPHOSPHORYLATION/CT  
 E E3+ALL  
 E E2+ALL  
 L9 5081 "DEPHOSPHORYLATION, BIOLOGICAL"/CT  
 E DRUG SCREENING/CT  
 E E3+ALL  
 E CHEMICAL LIBRARY/CT  
 E E3+ALL  
 E E5+ALL  
 E DRUG DISCOVERY/CT  
 E E3+ALL  
 L10 39357 DRUG DISCOVERY+NT/CT  
 L11 12 L7-8 AND L9 AND L10  
 E SCHAEFFER E/AU  
 L12 48 E3, E16  
 L13 10984 PFIZER/CS, PA  
 L14 1 L11 AND L12  
 L15 1 L11 AND L13  
 L16 1 L14-15  
 L17 11 L11 NOT L16  
 L18 8 L17 AND (PY<=2000 OR AY<=2000 OR PRY<=2000 OR PD<20001031 OR AD

FILE 'REGISTRY' ENTERED AT 10:42:04 ON 30 SEP 2004  
 E MAPK-P/CN  
 E P-NITROPHENYLPHOSPHATE/CN  
 L19 1 E4

FILE 'HCAPLUS' ENTERED AT 10:44:39 ON 30 SEP 2004  
 E IMMUNOASSAY/CT  
 E E3+ALL  
 L20 51907 IMMUNOASSAY+OLD, NT/CT  
 E IMMUNOCHEMICAL ANALYSIS/CT  
 E E3+ALL  
 L21 3702 IMMUNOCHEMICAL ANALYSIS/CT (L) IMMUNOASSAY?  
 L22 3 L7-8 AND L9 AND L20-21  
 L23 0 L22 AND L12-13  
 L24 2 L22 AND (PY<=2000 OR AY<=2000 OR PRY<=2000 OR PD<20001031 OR AD  
 L25 10 L24 OR L18

FILE 'REGISTRY' ENTERED AT 10:48:06 ON 30 SEP 2004  
 E O-METHYLFLUOROPHOSPHATE/CN  
 L26 0 CH4FO3P  
 L27 85 C6H6NO6P AND C6/ES  
 L28 33 L27 NOT ((PMS OR MAN OR IDS)/CI OR COMPD OR COMPOUND OR UNSPECI  
 L29 25 L28 AND 4(1A) NITROPHENYL  
 L30 24 L29 NOT PHOSPHONIC (1A) ACID  
 L31 24 L19 OR L30

FILE 'HCAPLUS' ENTERED AT 10:57:29 ON 30 SEP 2004  
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 L33 4950 (PHOSPHORIC (1A) ACID (1A) MONO (1A) 4 (1A) NITROPHENYL (1A) ES  
 E PHOSPOPTEIN/CT  
 E PHOSPHOPROTEIN/CT  
 E PHOSPHOPROTEINS/CT  
 E E3+ALL  
 L34 123837 PHOSPHOPROTEINS+NT/CT  
 L35 3 L25 AND L32-34  
 E HIGH THROUGHPUT SCREENING/CT  
 E E3+ALL  
 L36 3930 HIGH THROUGHPUT SCREENING/CT  
 L37 19 L36 AND L7-8  
 L38 1 L37 AND L9  
 L39 0 L38 AND L12-13  
 L40 0 L38 AND (PY<=2000 OR AY<=2000 OR PRY<=2000 OR PD<20001031 OR AD

FILE 'EMBASE' ENTERED AT 11:27:02 ON 30 SEP 2004  
 L41 5352 L7-8  
 L42 73832 (E5.160.160.680. OR G1.550.100.690. OR H1.150.150.680.)/CT  
 L43 425108 (E1.50.150.225. OR E5.680.202. OR E1.800. OR E5.795.)/CT  
 L44 40 L41 AND L42 AND L43  
 E SCHAEFFER E/AU  
 L45 54 E3  
 L46 4201 PFIZER/CS  
 L47 0 L44 AND L45-46

L48 17 L44 AND PY<=2001  
 L49 45600 D4.680.695./CT  
 L50 3 L48 AND (L31 OR L33 OR L49)  
     SEL AN 2-4 6 9-14 L48  
     SEL AN 1 3 L50  
 L51 11 E1-12 AND (L48 OR L50)

FILE 'WPIX' ENTERED AT 12:49:49 ON 30 SEP 2004  
 L52 379 L8  
 L53 99105 (B11-C08? OR C11-C08? OR B11-C10? OR C11-C10?)/MC OR G01N033-53  
 L54 958 DEPHOSPHORYLAT?/BIX  
 L55 12 L52 AND L53 AND L54  
     E SCHAEFFER E/AU  
 L56 10 E3  
 L57 4980 PFIZER/CS, PA  
 L58 0 L55 AND L56-57  
 L59 4 L55 NOT (PY>2001 OR PRY>2001 OR AY>2001)  
     SEL AN 1-3  
 L60 3 E1-3 AND L59

=> b hcap  
 FILE 'HCAPLUS' ENTERED AT 13:00:55 ON 30 SEP 2004  
 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT.  
 PLEASE SEE "HELP USAGETERMS" FOR DETAILS.  
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FILE COVERS 1907 - 30 Sep 2004 VOL 141 ISS 14  
 FILE LAST UPDATED: 29 Sep 2004 (20040929/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

=> d all 116 tot

L16 ANSWER 1 OF 1 HCAPLUS COPYRIGHT 2004 ACS on STN  
 AN 2002:331864 HCAPLUS  
 DN 136:337033  
 ED Entered STN: 03 May 2002  
 TI Drug screening for inhibitors of mouse protein tyrosine phosphatase PTPbr7 and their use in regulating nerve growth factor  
 IN Schaeffer, Eric  
 PA Pfizer Products Inc., USA  
 SO Eur. Pat. Appl., 14 pp.  
 CODEN: EPXXDW  
 DT Patent  
 LA English  
 IC ICM C12Q001-42  
 CC 7-1 (Enzymes)  
 Section cross-reference(s): 1, 13  
 FAN.CNT 1

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI EP 1201764	A2	20020502	EP 2001-124850	20011018
PI EP 1201764	A3	20040107		
		R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR		
		US 2002086342	A1 20020704 US 2001-437	20011031
		JP 2002355066	A2 20021210 JP 2001-334245	20011031
PRAI US 2000-244539P	P	20001031		

CLASS  
 PATENT NO. CLASS PATENT FAMILY CLASSIFICATION CODES  
 EP 1201764 ICM C12Q001-42

EP 1201764 ECLA C12Q001/42

AB The present invention relates to novel assays for the identification of agents that inhibit the catalytic activity of mouse PTPbr7. Substrates for dephosphorylation include phosphorylated mitogen-activating kinase and small peptides contain phosphotyrosine residues. An assay for identifying inhibitors of PTPbr7 includes an assay buffer containing 50 mM TRIS, 0.15 M NaCl, 5 mM DTT, 0.1% BSA at pH 7.4 and a volume of 25.μL. The assay is terminated by adding malachite green dye, ammonium molybdate, and Tween-20 with incubation period of 15 min. Thereafter, the optical d. of free inorg. phosphate is spectrophotometrically measured at 620 nm and compared with a set of stds., containing varied amts. of inorg. phosphate. The invention also provides pharmaceutical compns. comprising such agents identified using the assays of the invention. The invention further provides methods of treatment comprising administering such pharmaceutical compns.

ST drug screening inhibitor PTPbr7; protein tyrosine phosphatase br7 human regulation nerve growth factor; fusion protein GST histidine tag PTPbr7

IT Signal transduction, biological (PTPbr7 as neg. regulator of nerve growth factor; drug screening for inhibitors of human protein tyrosine phosphatase PTPbr7 and their use in regulating nerve growth factor)

IT Chimeric gene  
RL: BSU (Biological study, unclassified); BUU (Biological use, unclassified); BIOL (Biological study); USES (Uses)  
(PTPbr7 catalytic domain and GST encoded by; drug screening for inhibitors of human protein tyrosine phosphatase PTPbr7 and their use in regulating nerve growth factor)

IT Chimeric gene  
RL: BSU (Biological study, unclassified); BUU (Biological use, unclassified); BIOL (Biological study); USES (Uses)  
(PTPbr7 catalytic domain and histidine (6) tag encoded by; drug screening for inhibitors of human protein tyrosine phosphatase PTPbr7 and their use in regulating nerve growth factor)

IT Axon  
(PTPbr7 inhibitor in stimulating growth of; drug screening for inhibitors of human protein tyrosine phosphatase PTPbr7 and their use in regulating nerve growth factor)

IT Brain  
(PTPbr7 mRNA in; drug screening for inhibitors of human protein tyrosine phosphatase PTPbr7 and their use in regulating nerve growth factor)

IT Drug screening  
Mus  
(drug screening for inhibitors of human protein tyrosine phosphatase PTPbr7 and their use in regulating nerve growth factor)

IT Dephosphorylation, biological  
(of phospho-MAPK or peptides containing phosphotyrosines by PTPbr7, inhibitors of; drug screening for inhibitors of human protein tyrosine phosphatase PTPbr7 and their use in regulating nerve growth factor)

IT Peptides, biological studies  
RL: BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
(phosphotyrosine, PTPbr7 in dephosphorylation of; drug screening for inhibitors of human protein tyrosine phosphatase PTPbr7 and their use in regulating nerve growth factor)

IT 9061-61-4, Nerve growth factor  
RL: BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
(PTPbr7 as neg. regulator of; drug screening for inhibitors of human protein tyrosine phosphatase PTPbr7 and their use in regulating nerve growth factor)

IT 142243-02-5, Mitogen activated protein kinase  
RL: BSU (Biological study, unclassified); BIOL (Biological study)  
(PTPbr7 in dephosphorylation of; drug screening for inhibitors of human protein tyrosine phosphatase PTPbr7 and their use in regulating nerve growth factor)

IT 373389-66-3, PTPBR7

RL: ADV (Adverse effect, including toxicity); BSU (Biological study, unclassified); BIOL (Biological study)  
 (PTPbr7, of mouse; drug screening for inhibitors of human protein tyrosine phosphatase PTPbr7 and their use in regulating nerve growth factor)

IT 148851-08-5 416848-49-2  
 RL: BSU (Biological study, unclassified); BUU (Biological use, unclassified); BIOL (Biological study); USES (Uses)  
 (amino acid sequence for PTPbr7 peptide substrate; drug screening for inhibitors of human protein tyrosine phosphatase PTPbr7 and their use in regulating nerve growth factor)

IT 13721-39-6, Sodium orthovanadate  
 RL: BUU (Biological use, unclassified); BIOL (Biological study); USES (Uses)  
 (as non-specific tyrosine phosphate inhibitor in drug screening assay; drug screening for inhibitors of human protein tyrosine phosphatase PTPbr7 and their use in regulating nerve growth factor)

IT 754-02-9  
 RL: BUU (Biological use, unclassified); BIOL (Biological study); USES (Uses)  
 (as substrate for PTPbr7 dephosphorylation in drug screening assay; drug screening for inhibitors of human protein tyrosine phosphatase PTPbr7 and their use in regulating nerve growth factor)

IT 50812-37-8D, Glutathione-S-transferase, PTPbr7 catalytic domain fusion product with  
 RL: BUU (Biological use, unclassified); BIOL (Biological study); USES (Uses)  
 (drug screening for inhibitors of human protein tyrosine phosphatase PTPbr7 and their use in regulating nerve growth factor)

IT 71-00-1D, L-Histidine, PTPbr7 catalytic domain fusion product with  
 RL: BUU (Biological use, unclassified); BIOL (Biological study); USES (Uses)  
 (tag; drug screening for inhibitors of human protein tyrosine phosphatase PTPbr7 and their use in regulating nerve growth factor)

IT 416933-31-8 416933-32-9  
 RL: PRP (Properties)  
 (unclaimed sequence; drug screening for inhibitors of mouse protein tyrosine phosphatase PTPbr7 and their use in regulating nerve growth factor)

=> d all 125 tot

L25 ANSWER 1 OF 10 HCPLUS COPYRIGHT 2004 ACS on STN  
 AN 2003:376284 HCPLUS  
 DN 138:363219  
 ED Entered STN: 16 May 2003  
 TI Methods for the screening of agents modulating protein tyrosine phosphatase L1 (PTPL1) binding to the NMDA receptor or PTPL1-mediated dephosphorylation of NMDA receptor for the treatment of human diseases  
 IN Jerecic, Jasna; Braithwaite, Steven; Kask, Kalle; Liu, Jenkuei; Melcher, Thorsten  
 PA USA  
 SO U.S. Pat. Appl. Publ., 16 pp., Cont.-in-part of U.S. Ser. No. 774,481.  
 CODEN: USXXCO  
 DT Patent  
 LA English  
 IC ICM A61K031-00  
 ICS C12Q001-68; G01N033-53; G01N033-567  
 NCL 435007200; 435006000; 514001000  
 CC 2-8 (Mammalian Hormones)  
 Section cross-reference(s): 63  
 FAN.CNT 3

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 2003092071	A1	20030515	US 2002-246837	20020918 <--
US 2001049348	A1	20011206	US 2001-774481	20010130 <--
US 6521414	B2	20030218		
US 2004072275	A1	20040415	US 2003-633109	20030801 <--

PRAI US 2000-179453P P 20000201 <--  
 US 2001-774481 A2 20010130  
 US 2002-246837 A2 20020918

## CLASS

PATENT NO.	CLASS	PATENT FAMILY CLASSIFICATION CODES
US 2003092071	ICM	A61K031-00 ICS C12Q001-68; G01N033-53; G01N033-567 NCL 435007200; 435006000; 514001000
US 2004072275	ECLA	C12Q001/42; G01N033/94B <--
AB	The present invention relates to the identification of a binding between NMDA receptor (NMDA-R) subunits and a protein tyrosine phosphatase (PTP), e.g., PTPL1. The present invention provides methods for screening a PTP agonist or antagonist that modulates NMDA-R signaling. The present invention also provides methods and compns. for treatment of disorders mediated by abnormal NMDA-R signaling. The present invention further provides methods for isolating PTPL1 from a biol. preparation	
ST	NMDA receptor protein tyrosine phosphatase PTPL1 screening treatment	
IT	Nervous system, disease (Huntington's chorea; methods for the screening of agents modulating protein tyrosine phosphatase L1 (PTPL1) binding to the NMDA receptor or PTPL1-mediated dephosphorylation of NMDA receptor for the treatment of human diseases)	
IT	Glutamate receptors RL: BSU (Biological study, unclassified); BIOL (Biological study) (NMDA-binding; methods for the screening of agents modulating protein tyrosine phosphatase L1 (PTPL1) binding to the NMDA receptor or PTPL1-mediated dephosphorylation of NMDA receptor for the treatment of human diseases)	
IT	Drug tolerance (alc.; methods for the screening of agents modulating protein tyrosine phosphatase L1 (PTPL1) binding to the NMDA receptor or PTPL1-mediated dephosphorylation of NMDA receptor for the treatment of human diseases)	
IT	Pain (chronic; methods for the screening of agents modulating protein tyrosine phosphatase L1 (PTPL1) binding to the NMDA receptor or PTPL1-mediated dephosphorylation of NMDA receptor for the treatment of human diseases)	
IT	Mental disorder (dementia; methods for the screening of agents modulating protein tyrosine phosphatase L1 (PTPL1) binding to the NMDA receptor or PTPL1-mediated dephosphorylation of NMDA receptor for the treatment of human diseases)	
IT	Brain, disease (injury; methods for the screening of agents modulating protein tyrosine phosphatase L1 (PTPL1) binding to the NMDA receptor or PTPL1-mediated dephosphorylation of NMDA receptor for the treatment of human diseases)	
IT	Alzheimer's disease Analgesics Anti-Alzheimer's agents Anticonvulsants Antipsychotics Dephosphorylation, biological Drug dependence Drug screening Epilepsy Human Molecular association Nervous system agents Schizophrenia (methods for the screening of agents modulating protein tyrosine phosphatase L1 (PTPL1) binding to the NMDA receptor or PTPL1-mediated dephosphorylation of NMDA receptor for the treatment of human diseases)	
IT	Nerve, disease (motor; methods for the screening of agents modulating protein tyrosine phosphatase L1 (PTPL1) binding to the NMDA receptor or PTPL1-mediated dephosphorylation of NMDA receptor for the treatment of human diseases)	
IT	Nerve, disease (neuropathy; methods for the screening of agents modulating protein tyrosine phosphatase L1 (PTPL1) binding to the NMDA receptor or PTPL1-mediated dephosphorylation of	

NMDA receptor for the treatment of human diseases)  
 IT Signal transduction, biological  
 (of activated NMDA receptor; methods for screening of agents modulating  
 protein tyrosine phosphatase L1 (PTPL1)  
 binding to the NMDA receptor or PTPL1-mediated dephosphorylation of  
 NMDA receptor for treatment of human diseases)  
 IT Mental disorder  
 (psychosis; methods for the screening of agents modulating  
 protein tyrosine phosphatase L1 (PTPL1)  
 binding to the NMDA receptor or PTPL1-mediated dephosphorylation of  
 NMDA receptor for the treatment of human diseases)  
 IT Nervous system, disease  
 (spinocerebellar degeneration; methods for the screening of agents  
 modulating protein tyrosine phosphatase  
 L1 (PTPL1) binding to the NMDA receptor or PTPL1-mediated  
 dephosphorylation of NMDA receptor for the treatment of human diseases)  
 IT Brain, disease  
 (stroke; methods for the screening of agents modulating protein  
 tyrosine phosphatase L1 (PTPL1) binding to the NMDA  
 receptor or PTPL1-mediated dephosphorylation of NMDA receptor for the  
 treatment of human diseases)  
 IT Head, disease  
 (trauma; methods for the screening of agents modulating protein  
 tyrosine phosphatase L1 (PTPL1) binding to the NMDA  
 receptor or PTPL1-mediated dephosphorylation of NMDA receptor for the  
 treatment of human diseases)  
 IT 300851-68-7, PTP-L1  
 RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL  
 (Biological study); USES (Uses)  
 (agonists and antagonists; methods for the screening of agents  
 modulating protein tyrosine phosphatase  
 L1 (PTPL1) binding to the NMDA receptor or PTPL1-mediated  
 dephosphorylation of NMDA receptor for the treatment of human diseases)

L25 ANSWER 2 OF 10 HCAPLUS COPYRIGHT 2004 ACS on STN  
 AN 2002:575291 HCAPLUS  
 DN 137:137209  
 ED Entered STN: 02 Aug 2002  
 TI Three dimensional format biochips  
 IN Fagnani, Roberto; Hahn, Sonnkap; Dong, Xiaofan; Pircher, Tony; Matsumoto,  
 Sandra; Tsinberg, Pavel  
 PA Biocept, Inc., USA  
 SO PCT Int. Appl., 45 pp.  
 CODEN: PIXXD2  
 DT Patent  
 LA English  
 IC ICM C12Q001-68  
 CC 9-1 (Biochemical Methods)

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	WO 2002059372	A2	20020801	WO 2001-US51265	20011026 <--
	WO 2002059372	A3	20020919		
				W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG	
	EP 1328810	A2	20030723	EP 2001-994529	20011026 <--
				R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR	
	JP 2004518138	T2	20040617	JP 2002-559854	20011026 <--
PRAI	US 2000-243699P	P	20001026. <--		
	WO 2001-US51265	W	20011026		

CLASS

	PATENT NO.	CLASS	PATENT FAMILY CLASSIFICATION CODES
	WO 2002059372	ICM	C12Q001-68
	JP 2004518138	FTERM	4B024/AA11; 4B024/AA19; 4B024/AA20; 4B024/CA01; 4B024/CA11; 4B024/HA11; 4B024/HA20; 4B029/AA07; 4B029/AA21; 4B029/BB15; 4B029/BB20; 4B029/CC01; 4B029/CC02; 4B029/CC05; 4B029/FA12; 4B029/FA15;

4B063/QA01; 4B063/QA05; 4B063/QQ42; 4B063/QQ49;  
 4B063/QQ52; 4B063/QQ79; 4B063/QR32; 4B063/QR35;  
 4B063/QR38; 4B063/QR48; 4B063/QR56; 4B063/QR82;  
 4B063/QS32; 4B063/QS36; 4B063/QX02 <--

AB A biochip is formed with a plurality of optically clear hydrogel cells attached to the top surface of a solid substrate in the form of an array. Each of the cells is formed of a hydrogel of polyethylene glycol, polypropylene glycol or a copolymer thereof having reactive isocyanate groups. Nonhybridization binding entities are immobilized in these cells, which entities are effective to selectively sequester a target protein or other comparable biomol. Different binding entities are immobilized in different cells to create a biochip that can be used to assay for a number of target biomols.

ST biochip hydrogel biomol detection

IT Genetic element  
 RL: BSU (Biological study, unclassified); BIOL (Biological study)  
 (ERE (estrogen-responsive element); three dimensional format biochips)

IT Antibodies and Immunoglobulins  
 RL: ARU (Analytical role, unclassified); ANST (Analytical study)  
 (IgG; three dimensional format biochips)

IT Prostate-specific antigen  
 RL: BSU (Biological study, unclassified); BIOL (Biological study)  
 (antibodies to; three dimensional format biochips)

IT Polyoxyalkylenes, uses  
 RL: DEV (Device component use); USES (Uses)  
 (copolymers; three dimensional format biochips)

IT DNA  
 RL: ANT (Analyte); BSU (Biological study, unclassified); ANST (Analytical study); BIOL (Biological study)  
 (double-stranded; three dimensional format biochips)

IT Immunoassay  
 (enzyme-linked immunosorbent assay; three dimensional format biochips)

IT Albumins, analysis  
 RL: ARU (Analytical role, unclassified); ANST (Analytical study)  
 (serum, bovine; three dimensional format biochips)

IT Brain  
 Conformation  
 Dephosphorylation, biological

Diffusion

Electric field

Human

Molecular association

Nucleic acid hybridization

Phosphorylation

Viscosity

pH  
 (three dimensional format biochips)

IT Antigens  
 Transferrins  
 RL: ANT (Analyte); ANST (Analytical study)  
 (three dimensional format biochips)

IT Ferritins  
 RL: ARU (Analytical role, unclassified); BSU (Biological study, unclassified); ANST (Analytical study); BIOL (Biological study)  
 (three dimensional format biochips)

IT Proteins  
 RL: ARU (Analytical role, unclassified); PEP (Physical, engineering or chemical process); PYP (Physical process); ANST (Analytical study); PROC (Process)  
 (three dimensional format biochips)

IT Calmodulins

Estrogen receptors  
 RL: BSU (Biological study, unclassified); BIOL (Biological study)  
 (three dimensional format biochips)

IT Chelates  
 RL: BUU (Biological use, unclassified); BIOL (Biological study); USES (Uses)  
 (three dimensional format biochips)

IT Enzymes, biological studies  
 RL: BUU (Biological use, unclassified); BIOL (Biological study); USES (Uses)  
 (three dimensional format biochips)

IT Peptides, biological studies  
 RL: BUU (Biological use, unclassified); BIOL (Biological study); USES (Uses)  
 (three dimensional format biochips)

IT Receptors  
 RL: BUU (Biological use, unclassified); BIOL (Biological study); USES (Uses)  
 (three dimensional format biochips)

IT Antibodies and Immunoglobulins  
 RL: PEP (Physical, engineering or chemical process); PYP (Physical process); PROC (Process)  
 (three dimensional format biochips)

IT Macroglobulins  
 RL: ANT (Analyte); BSU (Biological study, unclassified); ANST (Analytical study); BIOL (Biological study)  
 (.alpha.2; three dimensional format biochips)

IT 79747-53-8, Yersinia outer membrane protein 2b 81156-93-6 106436-17-3  
 117592-22-0 118447-68-0 127212-49-1 129785-85-9 149261-42-7  
 150268-17-0 151419-98-6 166664-90-0 176042-83-4 198754-34-6  
 300857-98-1, Leukocyte antigen related protein tyrosine phosphatase 444166-73-8 444166-74-9 444166-75-0  
 444166-76-1 444166-77-2 444166-78-3 444166-79-4  
 RL: ANT (Analyte); ANST (Analytical study)  
 (three dimensional format biochips)

IT 9002-07-7, Trypsin 444166-80-7  
 RL: ANT (Analyte); BSU (Biological study, unclassified); ANST (Analytical study); BIOL (Biological study)  
 (three dimensional format biochips)

IT 11128-99-7, Angiotensin II 361540-77-4, Calcineurin  
 RL: BSU (Biological study, unclassified); BIOL (Biological study)  
 (three dimensional format biochips)

IT 59828-41-0, HYPOL  
 RL: BUU (Biological use, unclassified); BIOL (Biological study); USES (Uses)  
 (three dimensional format biochips)

IT 124-38-9, Carbon dioxide, processes  
 RL: CPS (Chemical process); PEP (Physical, engineering or chemical process); PROC (Process)  
 (three dimensional format biochips)

IT 25322-68-3D, Polyethylene glycol, copolymers 25322-69-4D, Polypropylene glycol, copolymers 181057-68-1, HYPOL PreMA G-50  
 RL: DEV (Device component use); USES (Uses)  
 (three dimensional format biochips)

IT 444272-42-8 444272-43-9 444272-44-0  
 RL: PRP (Properties)  
 (unclaimed sequence; three dimensional format biochips)

L25 ANSWER 3 OF 10 HCPLUS COPYRIGHT 2004 ACS on STN  
 AN 2002:408826 HCPLUS  
 DN 137:5025  
 ED Entered STN: 31 May 2002  
 TI T cell protein tyrosine phosphatase  
 inhibitors and activators for development of treatments for hematologic malignancies and autoimmune diseases  
 IN McGlade, Jane C.; Simoncic, Paul Daniel; Tremblay, Michael  
 PA The Hospital for Sick Children, Can.; McGill University  
 SO PCT Int. Appl., 64 pp.  
 CODEN: PIXXD2  
 DT Patent  
 LA English  
 IC ICM C12Q001-42  
 ICS G01N033-566; G01N033-68; A61K038-47; A61P037-04; A61P037-06;  
 G01N033-573  
 CC 15-10 (Immunochemistry)  
 Section cross-reference(s): 1, 3, 63  
 FAN.CNT 1

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2002042489	A1	20020530	WO 2001-CA1679	20011127 <--
WO 2002042489	C2	20021031		
			W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG	
AU 2002020412	A5	20020603	AU 2002-20412	20011127 <--

PRAI CA 2000-2326952 A 20001127 <--  
 WO 2001-CA1679 W 20011127

CLASS

PATENT NO.	CLASS	PATENT FAMILY CLASSIFICATION CODES
WO 2002042489	ICM	C12Q001-42
	ICS	G01N033-566; G01N033-68; A61K038-47; A61P037-04; A61P037-06; G01N033-573

AB This invention relates to T cell **protein tyrosine phosphatase** (TCPTP) and more particularly to its role in cell signaling and interaction with the JAK family of tyrosine kinases. In particular, the invention involves the use of TCPTP for the development of treatments for malignancies and autoimmune conditions involving inappropriate JAK kinase signaling as well as for the identification of inhibitors and activators of this phosphatase. The invention may also be used to rule out TCPTP inhibition in selecting potential anti-diabetic and anti-obesity PTP1B inhibitors without immune suppression.

ST T cell **protein tyrosine phosphatase** hematol malignancy; autoimmune disease T cell **protein tyrosine phosphatase**; JAK kinase T cell signaling immunosuppressant

IT Antitumor agents  
 Autoimmune disease  
 Dephosphorylation, biological  
 Drug design  
 Drug screening  
 Hematopoietic precursor cell  
 Peptidomimetics  
 Phosphorylation, biological  
 Signal transduction, biological  
 T cell (lymphocyte)  
 Transplant and Transplantation  
 (T cell **protein tyrosine phosphatase** inhibitors and activators for development of treatments for hematol. malignancies and autoimmune diseases)

IT Fusion proteins (chimeric proteins)  
 RL: BSU (Biological study, unclassified); BIOL (Biological study)  
 (T cell **protein tyrosine phosphatase** inhibitors and activators for development of treatments for hematol. malignancies and autoimmune diseases)

IT Antisense oligonucleotides  
 RL: BSU (Biological study, unclassified); DGN (Diagnostic use); THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
 (T cell **protein tyrosine phosphatase** inhibitors and activators for development of treatments for hematol. malignancies and autoimmune diseases)

IT Gene, animal  
 Proteins  
 RL: ARG (Analytical reagent use); BSU (Biological study, unclassified); DGN (Diagnostic use); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); USES (Uses)  
 (TCPTP; T cell **protein tyrosine phosphatase** inhibitors and activators for development of treatments for hematol. malignancies and autoimmune diseases)

IT Nucleic acids  
 RL: BSU (Biological study, unclassified); DGN (Diagnostic use); THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
 (antisense; T cell **protein tyrosine phosphatase** inhibitors and activators for development of treatments for hematol. malignancies and autoimmune diseases)

IT Drug delivery systems  
 (carriers; T cell **protein tyrosine phosphatase** inhibitors and activators for development of treatments for hematol. malignancies and autoimmune diseases)

IT Lymphocyte  
 (disease, proliferation defect; T cell **protein tyrosine phosphatase** inhibitors and activators for development of treatments for hematol. malignancies and autoimmune diseases)

IT Hematopoiesis  
 (disorders; T cell **protein tyrosine phosphatase** inhibitors and activators for development of treatments for hematol. malignancies and autoimmune diseases)

IT Neoplasm  
 (hematol.; T cell **protein tyrosine phosphatase** inhibitors and activators for development of treatments for hematol. malignancies and autoimmune diseases)

IT Antidiabetic agents  
 Antiobesity agents  
 (immunosuppression examination; T cell protein tyrosine phosphatase inhibitors and activators for development of treatments for hematol. malignancies and autoimmune diseases)

IT Carbohydrates, analysis  
 RL: ARU (Analytical role, unclassified); ANST (Analytical study)  
 (metabolism; T cell protein tyrosine phosphatase inhibitors and activators for development of treatments for hematol. malignancies and autoimmune diseases)

IT Diabetes mellitus  
 (resistance; T cell protein tyrosine phosphatase inhibitors and activators for development of treatments for hematol. malignancies and autoimmune diseases)

IT Anti-inflammatory agents  
 Immunostimulants  
 Immunosuppressants  
 (screening; T cell protein tyrosine phosphatase inhibitors and activators for development of treatments for hematol. malignancies and autoimmune diseases)

IT Mutagenesis  
 (site-directed, deletion; T cell protein tyrosine phosphatase inhibitors and activators for development of treatments for hematol. malignancies and autoimmune diseases)

IT Mutagenesis  
 (site-directed; T cell protein tyrosine phosphatase inhibitors and activators for development of treatments for hematol. malignancies and autoimmune diseases)

IT 152478-56-3, JAK1 kinase 157482-36-5, JAK3 kinase 161384-16-3, JAK kinase  
 RL: ARU (Analytical role, unclassified); BSU (Biological study, unclassified); ANST (Analytical study); BIOL (Biological study)  
 (T cell protein tyrosine phosphatase inhibitors and activators for development of treatments for hematol. malignancies and autoimmune diseases)

IT 300842-01-7, T Cell Protein tyrosine phosphatase  
 RL: ANT (Analyte); ARU (Analytical role, unclassified); BSU (Biological study, unclassified); DGN (Diagnostic use); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); USES (Uses)  
 (inhibitor and activator; T cell protein tyrosine phosphatase inhibitors and activators for development of treatments for hematol. malignancies and autoimmune diseases)

IT 60-18-4, Tyrosine, biological studies  
 RL: BSU (Biological study, unclassified); BIOL (Biological study)  
 (phosphorylation; T cell protein tyrosine phosphatase inhibitors and activators for development of treatments for hematol. malignancies and autoimmune diseases)

IT 432838-28-3 432838-29-4 432838-30-7 432838-31-8 432838-32-9  
 432838-33-0 432838-35-2 432838-36-3 432838-37-4 432838-38-5  
 432838-39-6 432838-40-9 432838-41-0  
 RL: PRP (Properties)  
 (unclaimed sequence; t cell protein tyrosine phosphatase inhibitors and activators for development of treatments for hematol. malignancies and autoimmune diseases)

RE.CNT 16 THERE ARE 16 CITED REFERENCES AVAILABLE FOR THIS RECORD

RE

- (1) Df Jesus, I; WO 0036111 A 2000
- (2) Flint, A; PROCEEDINGS OF THE NATIONAL ACADEMY OF SCIENCES OF THE UNITED STATES 1997, V94(5), P1680 HCAPLUS
- (3) Gjorloff-Wingren, A; EUROPEAN JOURNAL OF IMMUNOLOGY 2000, V30(8), P2412 HCAPLUS
- (4) Hui, Z; WO 0075339 A 2000 HCAPLUS
- (5) Ibarra-Sanchez, M; SEMIN IMMUNOL 2000, V12(4), P379 HCAPLUS
- (6) Ihle, J; SEMIN IMMUNOLOGY 1995, V7, P247 HCAPLUS
- (7) Lp, H; WO 9936548 A 1999 HCAPLUS
- (8) Maegawa, H; BIOCHEMICAL AND BIOPHYSICAL RESEARCH COMMUNICATIONS 1996, V228(1), P122 HCAPLUS
- (9) Maegawa, H; JOURNAL OF BIOLOGICAL CHEMISTRY 1999, V274(42), P30236 HCAPLUS
- (10) Novo Nordisk As; WO 0117516 A 2001 HCAPLUS
- (11) Sugen Inc; WO 9827092 A 1998 HCAPLUS
- (12) Tiganis, T; JOURNAL OF BIOLOGICAL CHEMISTRY 1997, V272(34), P21548 HCAPLUS
- (13) Tiganis, T; MOLECULAR AND CELLULAR BIOLOGY 1998, V18(3), P1622 HCAPLUS
- (14) Univ Rockefeller; WO 9923493 A 1999 HCAPLUS
- (15) Wimmer, M; HISTOCHEMISTRY AND CELL BIOLOGY 1999, V111(2), P135 HCAPLUS
- (16) You-Ten, K; JOURNAL OF EXPERIMENTAL MEDICINE 1997, V186(5), P683 HCAPLUS

L25 ANSWER 4 OF 10 HCPLUS COPYRIGHT 2004 ACS on STN  
 AN 2001:618190 HCPLUS

DN 135:207447

ED Entered STN: 24 Aug 2001

TI Fluorescent assay for protein tyrosine phosphatases

IN Flint, Andrew J.; Cool, Deborah E.

PA Ceptyr, Inc., USA

SO PCT Int. Appl., 79 pp.

CODEN: PIXXD2

DT Patent

LA English

IC ICM C12Q001-00

CC 7-1 (Enzymes)

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	WO 2001061031	A2	20010823	WO 2001-US5180	20010213 <--
	WO 2001061031	A3	20020307		
	WO 2001061031	C2	20021017		
		W:	AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM		
		RW:	GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG		
	US 2002009762	A1	20020124	US 2001-788626	20010213 <--
	EP 1257824	A2	20021120	EP 2001-910900	20010213 <--
		R:	AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR		
PRAI	US 2000-181769P	P	20000214		
	WO 2001-US5180	W	20010213		

CLASS

PATENT NO.	CLASS	PATENT FAMILY CLASSIFICATION CODES
WO 2001061031	ICM	C12Q001-00

AB The invention relates in part to screening assays for identifying agents that alter the interaction between a protein tyrosine phosphatase (PTP) and its tyrosine phosphorylated polypeptide substrate, using fluorescence energy signals generated by detectably labeled substrates. Assays are provided in certain embodiments, including high throughput screening assays, wherein candidate agents are screened by fluorescence polarization for their ability to influence (i) binding of substrate trapping mutant PTPs to substrates, or (ii) dephosphorylation of tyrosine phosphorylated substrates by PTPs.

ST protein tyrosine phosphatase detn  
 fluorescence polarization

IT Phosphoproteins

RL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses)  
 (P210bcr-c-abl, phosphopeptide derived from; fluorescent assay for protein tyrosine phosphatases)

IT Protein motifs

(PH domain, polypeptide containing, reaction terminator; fluorescent assay for protein tyrosine phosphatases)

IT Protein motifs

(PTB domain, polypeptide containing, reaction terminator; fluorescent assay for protein tyrosine phosphatases)

IT Protein motifs

(PTB-PID domain, polypeptide containing, reaction terminator; fluorescent assay for protein tyrosine phosphatases)

IT Protein motifs

(SH2 domain, polypeptide containing, reaction terminator; fluorescent assay for protein tyrosine phosphatases)

IT Phosphoproteins

RL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses)  
 (SHC, phosphopeptide derived from; fluorescent assay for protein tyrosine phosphatases)

IT Proteins, specific or class

RL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses)  
 (VCP, phosphopeptide derived from; fluorescent assay for protein tyrosine phosphatases)

IT Enzyme functional sites

(active; fluorescent assay for protein tyrosine phosphatases)  
 IT Resonant energy transfer  
 (fluorescence; fluorescent assay for protein tyrosine phosphatases)  
 IT Dephosphorylation, biological  
 Fluorescence  
 Fluorescent substances  
 Polarized fluorescence  
 (fluorescent assay for protein tyrosine phosphatases)  
 IT CD45 (antigen)  
 RL: ANT (Analyte); ANST (Analytical study)  
 (fluorescent assay for protein tyrosine phosphatases)  
 IT Phosphopeptides  
 RL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses)  
 (fluorescent assay for protein tyrosine phosphatases)  
 IT Drug screening  
 (fluorescent assay for protein tyrosine phosphatases in relation to)  
 IT Proteins, specific or class  
 RL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses)  
 (p130, p130cas, phosphopeptide derived from; fluorescent assay for protein tyrosine phosphatases)  
 IT Epidermal growth factor receptors  
 Insulin receptors  
 RL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses)  
 (phosphopeptide derived from; fluorescent assay for protein tyrosine phosphatases)  
 IT Phosphoproteins  
 RL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses)  
 (pp56lck, phosphopeptide derived from; fluorescent assay for protein tyrosine phosphatases)  
 IT Antibodies  
 RL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses)  
 (reaction terminator; fluorescent assay for protein tyrosine phosphatases)  
 IT Mutation  
 (substitution, of protein tyrosine phosphatase; fluorescent assay for protein tyrosine phosphatases)  
 IT TCR (T cell receptors)  
 RL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses)  
 (zeta chain, phosphopeptide derived from; fluorescent assay for protein tyrosine phosphatases)  
 IT 60-18-4, L-Tyrosine, biological studies  
 RL: BSU (Biological study, unclassified); BIOL (Biological study)  
 (-676, phosphatase wild-type Tyr replaced by; fluorescent assay for protein tyrosine phosphatases)  
 IT 247144-99-6, Alexa Fluor 488  
 RL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses)  
 (Alexa Fluor 488, fluorophore; fluorescent assay for protein tyrosine phosphatases)  
 IT 247145-86-4, Alexa Fluor 594  
 RL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses)  
 (Alexa Fluor 594, fluorophore; fluorescent assay for protein tyrosine phosphatases)  
 IT 146368-14-1  
 RL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses)  
 (Cy5, fluorophore; fluorescent assay for protein tyrosine phosphatases)  
 IT 330956-25-7, Leukocyte protein tyrosine phosphatase  
 RL: ANT (Analyte); ANST (Analytical study)  
 (LC-PTP, reaction terminator; fluorescent assay for protein tyrosine phosphatases)  
 IT 79747-53-8, Protein tyrosine phosphatase  
 196717-98-3, protein tyrosine phosphatase  
 PEST 300830-54-0, PTP-MEG1 300842-01-7, TC-PTP 300857-98-1,  
 Protein tyrosine phosphatase LAR  
 300865-11-6, protein tyrosine phosphatase 1B  
 300865-18-3, Protein tyrosine phosphatase  
 .mu. 300865-46-7, Protein tyrosine phosphatase .gamma. 301156-53-6, Protein

tyrosine phosphatase SHP2 301162-72-1, Protein  
 tyrosine phosphatase H1 306298-47-5, MAPK phosphatase  
 1  
 RL: ANT (Analyte); ANST (Analytical study)  
 (fluorescent assay for protein tyrosine  
 phosphatases)  
 IT 76823-03-5DP, 5-Carboxyfluorescein, reaction products with phosphopeptides  
 207554-36-7DP, reaction products with fluorescein 357164-39-7DP,  
 reaction products with fluorescein 357164-40-0DP, reaction products with  
 fluorescein  
 RL: ARG (Analytical reagent use); SPN (Synthetic preparation); ANST  
 (Analytical study); PREP (Preparation); USES (Uses)  
 (fluorescent assay for protein tyrosine  
 phosphatases)  
 IT 2321-07-5, Fluorescein 13558-31-1 82354-19-6, Texas Red 165599-63-3,  
 BODIPY-FL  
 RL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses)  
 (fluorophore; fluorescent assay for protein tyrosine  
 phosphatases)  
 IT 52-90-4, L-Cysteine, analysis 56-40-6, Glycine, analysis 56-41-7,  
 L-Alanine, analysis 56-84-8, L-Aspartic acid, analysis 56-85-9,  
 L-Glutamine, analysis 56-86-0, L-Glutamic acid, analysis 56-87-1,  
 L-Lysine, analysis 61-90-5, L-Leucine, analysis 63-68-3, L-Methionine,  
 analysis 63-91-2, L-Phenylalanine, analysis 70-47-3, L-Asparagine,  
 analysis 71-00-1, L-Histidine, analysis 72-18-4, L-Valine, analysis  
 73-22-3, L-Tryptophane, analysis 73-32-5, L-Isoleucine, analysis  
 74-79-3, L-Arginine, analysis 147-85-3, L-Proline, analysis  
 RL: ARU (Analytical role, unclassified); ANST (Analytical study)  
 (phosphatase wild-type Tyr replaced by; fluorescent assay for  
 protein tyrosine phosphatases)  
 IT 114051-78-4, Lck tyrosine kinase  
 RL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses)  
 (phosphopeptide derived from; fluorescent assay for protein  
 tyrosine phosphatases)  
 IT 300854-55-1, PTP-DEP1 300859-91-0, PTP-CD45 356771-02-3 356771-72-7  
 356771-80-7  
 RL: ANT (Analyte); ANST (Analytical study)  
 (reaction terminator; fluorescent assay for protein  
 tyrosine phosphatases)  
 IT 37353-31-4, vanadate  
 RL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses)  
 (reaction terminator; fluorescent assay for protein  
 tyrosine phosphatases)  
 IT 142243-02-5, MAP kinase  
 RL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses)  
 (substrate phosphopeptide derived from; fluorescent assay for  
 protein tyrosine phosphatases)  
 IT 313285-28-8 313285-34-6 313285-39-1 357464-46-1 357464-47-2  
 357464-48-3 357464-49-4 357464-50-7 357464-51-8 357464-52-9  
 357464-53-0 357464-54-1 357464-55-2 357464-56-3 357464-57-4  
 357464-58-5 357464-59-6 357464-60-9 357464-61-0 357464-62-1  
 357464-63-2 357464-64-3 357464-65-4 357464-66-5 357464-67-6  
 357464-68-7 357464-69-8 357464-70-1 357464-71-2 357464-72-3  
 357464-73-4 357464-74-5 357464-75-6 357464-76-7 357464-77-8  
 RL: PRP (Properties)  
 (unclaimed sequence; fluorescent assay for protein  
 tyrosine phosphatases)

L25 ANSWER 5 OF 10 HCAPLUS COPYRIGHT 2004 ACS on STN  
 AN 2001:582074 HCAPLUS  
 DN 135:163001  
 ED Entered STN: 10 Aug 2001  
 TI Interaction of NMDA receptor with protein tyrosine  
 phosphatase, screening for agents which modulate NMDA receptor  
 signaling, and therapeutic applications  
 IN Melcher, Thorsten; Kask, Kalev  
 PA Agy Therapeutics, Inc., USA  
 SO PCT Int. Appl., 34 pp.  
 CODEN: PIXXD2  
 DT Patent  
 LA English  
 IC ICM C12Q001-00  
 CC 2-8 (Mammalian Hormones)  
 Section cross-reference(s): 1, 7, 63  
 FAN.CNT 3

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
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PI	WO 2001057240	A2	20010809	WO 2001-US3049	20010130 <--
	W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM	RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG			
	EP 1255854	A2	20021113	EP 2001-908752	20010130 <--
	R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR				
	JP 2003521247	T2	20030715	JP 2001-555863	20010130 <--
PRAI	US 2000-179453P	P	20000201		
	WO 2001-US3049	W	20010130		

## CLASS

PATENT NO.	CLASS	PATENT FAMILY CLASSIFICATION CODES
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WO 2001057240	ICM	C12Q001-00
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AB The present invention relates to the identification of a binding between NMDA receptor (NMDA-R) subunits and a **protein tyrosine phosphatase** (PTP), e.g., PTPL1. The present invention provides methods for screening a PTP agonist or antagonist that modulates NMDA-R signaling. The present invention also provides methods and compns. for treatment of disorders mediated by abnormal NMDA-R signaling. The present invention further provides methods for isolating PTPL1 from a biol. preparation

ST NMDA receptor protein tyrosine phosphatase  
PTPL1 therapeutic; drug screening NMDA receptor protein tyrosine phosphatase PTPL1

IT Nervous system  
(Huntington's chorea, treatment of; interaction of NMDA receptor with **protein tyrosine phosphatase** PTPL1, screening for agents which modulate NMDA receptor signaling, and therapeutic applications)

IT Gene, animal  
RL: ANT (Analyte); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); USES (Uses)  
(NMDA receptor signaling modulator-encoding; interaction of NMDA receptor with **protein tyrosine phosphatase** PTPL1, screening for agents which modulate NMDA receptor signaling, and therapeutic applications)

IT Glutamate receptors  
RL: BPR (Biological process); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); PROC (Process); USES (Uses)  
(NMDA-binding; interaction of NMDA receptor with **protein tyrosine phosphatase** PTPL1, screening for agents which modulate NMDA receptor signaling, and therapeutic applications)

IT Mental disorder  
(dementia, treatment of; interaction of NMDA receptor with **protein tyrosine phosphatase** PTPL1, screening for agents which modulate NMDA receptor signaling, and therapeutic applications)

IT Analgesics  
(for chronic pain treatment; interaction of NMDA receptor with **protein tyrosine phosphatase** PTPL1, screening for agents which modulate NMDA receptor signaling, and therapeutic applications)

IT Brain, disease  
(injury, treatment of; interaction of NMDA receptor with **protein tyrosine phosphatase** PTPL1, screening for agents which modulate NMDA receptor signaling, and therapeutic applications)

IT Anti-Alzheimer's agents  
Anticonvulsants  
Antipsychotics  
Dephosphorylation, biological  
Drug screening  
Drugs  
Molecular association  
Signal transduction, biological  
(interaction of NMDA receptor with **protein tyrosine phosphatase** PTPL1, screening for agents which modulate NMDA receptor signaling, and therapeutic applications)

IT Nerve, disease

(motor, treatment of; interaction of NMDA receptor with protein tyrosine phosphatase PTPL1, screening for agents which modulate NMDA receptor signaling, and therapeutic applications)

IT Nerve, disease  
(neuropathy, treatment of; interaction of NMDA receptor with protein tyrosine phosphatase PTPL1, screening for agents which modulate NMDA receptor signaling, and therapeutic applications)

IT Phosphorylation, biological  
(protein; interaction of NMDA receptor with protein tyrosine phosphatase PTPL1, screening for agents which modulate NMDA receptor signaling, and therapeutic applications)

IT Nervous system  
(spinocerebellar degeneration, treatment of; interaction of NMDA receptor with protein tyrosine phosphatase PTPL1, screening for agents which modulate NMDA receptor signaling, and therapeutic applications)

IT Brain, disease  
(stroke, treatment of; interaction of NMDA receptor with protein tyrosine phosphatase PTPL1, screening for agents which modulate NMDA receptor signaling, and therapeutic applications)

IT Head  
(trauma, treatment of; interaction of NMDA receptor with protein tyrosine phosphatase PTPL1, screening for agents which modulate NMDA receptor signaling, and therapeutic applications)

IT Alcoholism

Drug dependence

Schizophrenia  
(treatment of; interaction of NMDA receptor with protein tyrosine phosphatase PTPL1, screening for agents which modulate NMDA receptor signaling, and therapeutic applications)

IT 79747-53-8P  
RL: BAC (Biological activity or effector, except adverse); BPR (Biological process); BSU (Biological study, unclassified); PUR (Purification or recovery); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); PROC (Process); USES (Uses)  
(L1 isoenzyme; interaction of NMDA receptor with protein tyrosine phosphatase PTPL1, screening for agents which modulate NMDA receptor signaling, and therapeutic applications)

L25 ANSWER 6 OF 10 HCAPLUS COPYRIGHT 2004 ACS on STN  
AN 2000:251041 HCAPLUS  
DN 133:70565  
ED Entered STN: 19 Apr 2000  
TI Structure-based design of a low molecular weight, nonphosphorus, nonpeptide, and highly selective inhibitor of protein-tyrosine phosphatase 1B  
AU Iversen, Lars Fogh; Andersen, Henrik Sune; Branner, Sven; Mortensen, Steen B.; Peters, Gunther H.; Norris, Kjeld; Olsen, Ole Hvilsted; Jeppesen, Claus Bekker; Lundt, Behrend F.; Ripka, William; Moller, Karin Bach; Moller, Niels Peter Hundahl  
CS Protein Chemistry, Bagsvaerd, DK-2880, Den.  
SO Journal of Biological Chemistry (2000), 275(14), 10300-10307  
CODEN: JBCHA3; ISSN: 0021-9258  
PB American Society for Biochemistry and Molecular Biology  
DT Journal  
LA English  
CC 7-3 (Enzymes)  
Section cross-reference(s): 1, 75  
AB Several protein-tyrosine phosphatases (PTPs) have been proposed to act as neg. regulators of insulin signaling. Recent studies have shown increased insulin sensitivity and resistance to obesity in PTP1B knockout mice, thus pointing to this enzyme as a potential drug target in diabetes. Structure-based design, guided by PTP mutants and x-ray protein crystallog., was used to optimize a relatively weak, nonphosphorus, nonpeptide general PTP inhibitor (2-(oxalyl-amino)-benzoic acid) into a highly selective PTP1B inhibitor. This was achieved by addressing residue 48 as a selectivity determining residue. By introducing a basic nitrogen in the core structure of the inhibitor, a salt bridge was formed to Asp-48 in PTP1B. In contrast, the basic nitrogen causes repulsion in other PTPs containing an asparagine in the equivalent position resulting in a remarkable selectivity for PTP1B. Importantly, this was accomplished while retaining the mol. weight of the inhibitor below 300 g/mol.

ST protein tyrosine phosphatase 1B inhibitor  
oxalylaminobenzoate; crystal structure protein tyrosine phosphatase 1B

IT Enzyme functional sites  
(active; structure-based design of a low mol. weight, nonphosphorus, nonpeptide, and highly selective inhibitor of protein-tyrosine phosphatase 1B)

IT Structure-activity relationship  
(enzyme-inhibiting; structure-based design of a low mol. weight, nonphosphorus, nonpeptide, and highly selective inhibitor of protein-tyrosine phosphatase 1B)

IT Enzyme kinetics  
(of inhibition; structure-based design of a low mol. weight, nonphosphorus, nonpeptide, and highly selective inhibitor of protein-tyrosine phosphatase 1B)

IT Conformation  
(protein; structure-based design of a low mol. weight, nonphosphorus, nonpeptide, and highly selective inhibitor of protein-tyrosine phosphatase 1B)

IT Conformational transition

Crystal structure  
Dephosphorylation, biological  
Drug design  
(structure-based design of a low mol. weight, nonphosphorus, nonpeptide, and highly selective inhibitor of protein-tyrosine phosphatase 1B)

IT 79747-53-8  
RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process)  
(1B; structure-based design of a low mol. weight, nonphosphorus, nonpeptide, and highly selective inhibitor of protein-tyrosine phosphatase 1B)

IT 79747-53-8D, complexes with oxalylaminobenzoate-based derivs.  
RL: PRP (Properties)  
(1B; structure-based design of a low mol. weight, nonphosphorus, nonpeptide, and highly selective inhibitor of protein-tyrosine phosphatase 1B)

IT 5651-01-4 243966-03-2 243967-41-1 243967-42-2 243985-35-5  
243985-58-2  
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); PRP (Properties); BIOL (Biological study)  
(structure-based design of a low mol. weight, nonphosphorus, nonpeptide, and highly selective inhibitor of protein-tyrosine phosphatase 1B)

IT 243967-41-1D, complexes with protein-tyrosine phosphatase 1B 243967-42-2D, complexes with protein-tyrosine phosphatase 1B  
RL: PRP (Properties)  
(structure-based design of a low mol. weight, nonphosphorus, nonpeptide, and highly selective inhibitor of protein-tyrosine phosphatase 1B)

RE.CNT 43 THERE ARE 43 CITED REFERENCES AVAILABLE FOR THIS RECORD

RE

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L25 ANSWER 7 OF 10 HCPLUS COPYRIGHT 2004 ACS on STN  
 AN 2000:185782 HCPLUS  
 DN 132:344765  
 ED Entered STN: 23 Mar 2000  
 TI 2-(Oxylamino)-benzoic acid is a general, competitive inhibitor of protein-tyrosine phosphatases  
 AU Andersen, Henrik Sune; Iversen, Lars Fogh; Jeppesen, Claus Bekker; Branner, Sven; Norris, Kjeld; Rasmussen, Hanne B.; Moller, Karin Bach; Moller, Niels Peter Hundahl  
 CS MedChem Research I, Bagsvaerd, DK-2880, Den.  
 SO Journal of Biological Chemistry (2000), 275(10), 7101-7108  
 CODEN: JBCHA3; ISSN: 0021-9258  
 PB American Society for Biochemistry and Molecular Biology  
 DT Journal  
 LA English  
 CC 7-3 (Enzymes)  
 Section cross-reference(s): 1, 75  
 AB Protein-tyrosine phosphatases (PTPs) are critically involved in regulation of signal transduction processes. Members of this class of enzymes are considered attractive therapeutic targets in several disease states, e.g. diabetes, cancer, and inflammation. However, most reported PTP inhibitors have been phosphorus-containing compds., tight binding inhibitors, and/or inhibitors that covalently modify the enzymes. We therefore embarked on identifying a general, reversible, competitive PTP inhibitor that could be used as a common scaffold for lead optimization for specific PTPs. We here report the identification of 2-(oxylamino)-benzoic acid (OBA) as a classical competitive inhibitor of several PTPs. X-ray crystallog. of PTP1B complexed with OBA and related non-phosphate low mol. weight derivs. reveals that the binding mode of these mols. to a large extent mimics that of the natural substrate including hydrogen bonding to the PTP signature motif. In addition, binding of OBA to the active site of PTP1B creates a unique arrangement involving Asp181, Lys120, and Tyr46. PTP inhibitors are essential tools in elucidating the biol. function of specific PTPs and they may eventually be developed into selective drug candidates. The unique enzyme kinetic features and the low mol. weight of OBA makes it an ideal starting point for further optimization.  
 ST protein tyrosine phosphatase inhibitor  
 oxalylaminobenzoate drug design; crystal structure protein tyrosine phosphatase inhibitor  
 IT Conformation  
 Conformational transition  
 Crystal structure  
 Dephosphorylation, biological  
 Drug design  
 Ionization  
 (2-(oxylamino)-benzoic acid is a general, competitive inhibitor of protein-tyrosine phosphatases)  
 IT Enzyme functional sites  
 (active; 2-(oxylamino)-benzoic acid is a general, competitive inhibitor of protein-tyrosine phosphatases)  
 IT Enzyme kinetics  
 (of inhibition; 2-(oxylamino)-benzoic acid is a general, competitive

inhibitor of protein-tyrosine phosphatases  
)

IT 5651-01-4, 2-(Oxalylamino)-benzoic acid 243967-43-3 243967-44-4  
243989-49-3 243989-50-6  
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); BIOL (Biological study)  
(2-(oxalylamino)-benzoic acid is a general, competitive inhibitor of protein-tyrosine phosphatases)

IT 5651-01-4D, 2-(Oxalylamino)-benzoic acid, complexes with protein-tyrosine phosphatases 79747-53-8D, Protein tyrosine phosphatase, complexes with oxalylaminobenzoate and derivs. 243967-44-4D, complexes with protein-tyrosine phosphatases 243989-49-3D, complexes with protein-tyrosine phosphatases 243989-50-6D, complexes with protein-tyrosine phosphatases  
RL: PRP (Properties)  
(2-(oxalylamino)-benzoic acid is a general, competitive inhibitor of protein-tyrosine phosphatases)

IT 79747-53-8, Protein tyrosine phosphatase  
RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process)  
(multiple forms of; 2-(oxalylamino)-benzoic acid is a general, competitive inhibitor of protein-tyrosine phosphatases)

RE.CNT 44 THERE ARE 44 CITED REFERENCES AVAILABLE FOR THIS RECORD

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L25 ANSWER 8 OF 10 HCPLUS COPYRIGHT 2004 ACS on STN

AN 2000:101696 HCPLUS

DN 132:304951

ED Entered STN: 13 Feb 2000

TI 3,6-Fluorescein diphosphate: a sensitive fluorogenic and chromogenic substrate for protein tyrosine phosphatases  
 AU Huang, Zheng; Wang, Qingping; Ly, Hoa D.; Govindarajan, Arvind; Scheigetz, John; Zamboni, Robert; Desmarais, Sylvie; Ramachandran, Chidambaram  
 CS Merck Frosst Center for Therapeutic Research, Dorval, QC, Can.  
 SO Journal of Biomolecular Screening (1999), 4 (6), 327-334  
 CODEN: JBISF3; ISSN: 1087-0571  
 PB Mary Ann Liebert, Inc.  
 DT Journal  
 LA English  
 CC 7-1 (Enzymes)  
 Section cross-reference(s): 1  
 AB A highly sensitive and continuous protein tyrosine phosphatase (PTPase) assay using 3,6-fluorescein diphosphate (FDP) is described. Leukocyte phosphatase CD45 (leukocyte common antigen), protein tyrosine phosphatase-1B, and leukocyte common antigen-related protein LAR preferentially hydrolyze FDP to fluorescein monophosphate (FMP) with Vmax and Km values comparable with those of phosphotyrosine peptide substrates. Further hydrolysis of FMP to fluorescein was less efficient because of increased Km values compared with those of FDP. FMP absorbs strongly at 445 nm and fluoresces intensely near 515 nm, both of which are insensitive to pH perturbations above pH 6. Its high catalytic efficiency, coupled with the highly sensitive dual detection in the visible wavelength region and wider pH operating range, make FDP the substrate of choice for PTPase inhibitor screening in HTS format and assay miniaturization.  
 ST protein tyrosine phosphatase substrate  
 fluorescein diphosphate  
 IT Dephosphorylation, biological  
     Drug screening  
     Fluorescence  
     Michaelis constant  
     (3,6-fluorescein diphosphate is a sensitive fluorogenic and chromogenic substrate for protein tyrosine phosphatases  
     )  
 IT 79747-53-8, Protein tyrosine phosphatase  
 RL: ANT (Analyte); ANST (Analytical study)  
     (3,6-fluorescein diphosphate is a sensitive fluorogenic and chromogenic substrate for protein tyrosine phosphatases  
     )  
 IT 134869-03-7, 3,6-Fluorescein diphosphate  
 RL: ARG (Analytical reagent use); BPR (Biological process); BSU (Biological study, unclassified); PRP (Properties); ANST (Analytical study); BIOL (Biological study); PROC (Process); USES (Uses)  
     (3,6-fluorescein diphosphate is a sensitive fluorogenic and chromogenic substrate for protein tyrosine phosphatases  
     )  
 IT 185252-56-6  
 RL: BPR (Biological process); BSU (Biological study, unclassified); PRP (Properties); BIOL (Biological study); PROC (Process)  
     (3,6-fluorescein diphosphate is a sensitive fluorogenic and chromogenic substrate for protein tyrosine phosphatases  
     )  
 RE.CNT 26 THERE ARE 26 CITED REFERENCES AVAILABLE FOR THIS RECORD  
 RE  
 (1) Burke, T; Biochem Biophys Res Commun 1994, V204, P129 HCPLUS  
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L25 ANSWER 9 OF 10 HCAPLUS COPYRIGHT 2004 ACS on STN  
 AN 1999:356398 HCAPLUS  
 DN 131:196127  
 ED Entered STN: 10 Jun 1999  
 TI Microtiter Well Assays for Protein Tyrosine Phosphatase Activities Directed against Phosphorylated Insulin Receptor or Insulin Receptor Substrate-1  
 AU Krutzfeldt, Jan; Grunweller, Arnold; Raasch, Walter; Drenckhan, Maren; Klein, Harald H.  
 CS Department of Internal Medicine 1, Medical University of Lubeck, Lubeck, D-23538, Germany  
 SO Analytical Biochemistry (1999), 271(1), 97-99  
 CODEN: ANBCA2; ISSN: 0003-2697  
 PB Academic Press  
 DT Journal  
 LA English  
 CC 7-1 (Enzymes)  
 AB A microwell-based assay system was developed that allows one to specifically measure protein tyrosine phosphatase (PTPase) activities directed against two proteins involved in insulin signaling. It represents a useful tool for the investigation of potential alterations in PTPase activities in different states of insulin resistance. Moreover, similar assays can be established for other membrane-bound and cytosolic tyrosine-phosphorylated proteins.  
 (c) 1999 Academic Press.  
 ST microtiter assay protein tyrosine phosphatase insulin receptor substrate  
 IT Phosphoproteins  
 RL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses)  
 (IRS-1 (insulin receptor substrate 1), phosphorylated, immobilized; microtiter well assays for protein tyrosine phosphatase activities directed against phosphorylated insulin receptor or insulin receptor substrate-1)  
 IT Antibodies  
 RL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses)  
 (biotinylated, against phosphotyrosine; microtiter well assays for protein tyrosine phosphatase activities directed against phosphorylated insulin receptor or insulin receptor substrate-1)  
 IT Immunoassay  
 (enzyme-linked immunosorbent assay; microtiter well assays for protein tyrosine phosphatase activities directed against phosphorylated insulin receptor or insulin receptor substrate-1)  
 IT Dephosphorylation, biological  
 (microtiter well assays for protein tyrosine phosphatase activities directed against phosphorylated insulin receptor or insulin receptor substrate-1)  
 IT Antibodies  
 RL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses)  
 (monoclonal, against insulin receptor or insulin receptor substrate-1; microtiter well assays for protein tyrosine phosphatase activities directed against phosphorylated insulin receptor or insulin receptor substrate-1)  
 IT Insulin receptors  
 RL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses)  
 (phosphorylated, immobilized; microtiter well assays for protein tyrosine phosphatase activities directed against phosphorylated insulin receptor or insulin receptor substrate-1)  
 IT 79747-53-8, Protein tyrosine phosphatase  
 RL: ANT (Analyte); ANST (Analytical study)  
 (microtiter well assays for protein tyrosine phosphatase activities directed against phosphorylated insulin receptor or insulin receptor substrate-1)  
 IT 9003-99-0D, Peroxidase, conjugates with streptavidin 9013-20-1D, Streptavidin, conjugates with horseradish peroxidase 28752-68-3, ABTS  
 RL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses)  
 (microtiter well assays for protein tyrosine

phosphatase activities directed against phosphorylated insulin receptor or insulin receptor substrate-1)

RE.CNT 13 THERE ARE 13 CITED REFERENCES AVAILABLE FOR THIS RECORD

RE

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L25 ANSWER 10 OF 10 HCAPLUS COPYRIGHT 2004 ACS on STN

AN 1998:550508 HCAPLUS

DN 129:187420

ED Entered STN: 31 Aug 1998

TI Tyrosine phosphorylated proteins (PSTPIP) found in the cleavage furrow that are substrates for PEST protein tyrosine phosphatases

IN Lasky, Laurence A.; Dowbenko, Donald J.

PA Genentech, Inc., USA

SO PCT Int. Appl., 113 pp.

CODEN: PIXXD2

DT Patent

LA English

IC ICM C12N015-12

ICS C07K014-47; G01N033-50; C07K016-18; C12N005-20

CC 13-3 (Mammalian Biochemistry)

Section cross-reference(s): 6

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	WO 9835037	A1	19980813	WO 1998-US1774	19980130 <--
	W: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, FI, GB, GE, GH, GM, GW, HU, ID, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM				
	RW: GH, GM, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG				
	ZA 9800651	A	19990727	ZA 1998-651	19980127 <--
	AU 9866493	A1	19980826	AU 1998-66493	19980130 <--
	AU 731570	B2	20010405		
	EP 980426	A1	20000223	EP 1998-908458	19980130 <--
	R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, FI				
	JP 2001503994	T2	20010327	JP 1998-534800	19980130 <--
	JP 3503951	B2	20040308		
	MX 9907114	A	20000228	MX 1999-7114	19990730 <--
PRAI	US 1997-798419	A	19970207	<--	
	US 1997-938829	A2	19970929	<--	
	WO 1998-US1774	W	19980130	<--	

CLASS

	PATENT NO.	CLASS	PATENT FAMILY CLASSIFICATION CODES
	WO 9835037	ICM	C12N015-12
		ICS	C07K014-47; G01N033-50; C07K016-18; C12N005-20
	WO 9835037	ECLA	C07K014/47 <--

AB Novel proteins that are substrates for dephosphorylation by the PEST family of protein tyrosine phosphatases and that are associated with the cleavage furrow are described and cDNAs encoding them are cloned from mouse. The protein appears to play a role in the polymerization of actin and so may be a target for the control of the process. The protein was identified as a ligand for PTP PEST in a yeast two-hybrid system using a cDNA bank from mouse Baf3 cells. The protein was found in actin-rich sites within the cell, specifically with the cortical actin cytoskeleton.

ST PSTPIP actin polymn cytokinesis; tyrosine dephosphorylation PSTPIP PTP PEST; cDNA PSTPIP mouse; cleavage furrow PSTPIP

IT Cytoskeleton  
 (PSTPIP association with cortical actin of; tyrosine phosphorylated proteins (PSTPIPs) found in cleavage furrow that are substrates for PEST protein tyrosine phosphatases)

IT Actins  
 RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process)  
 (PSTPIP interaction with; tyrosine phosphorylated proteins (PSTPIPs) found in cleavage furrow that are substrates for PEST protein tyrosine phosphatases)

IT Embryo, animal  
 (PSTPIP levels in, as function of developmental stage; tyrosine phosphorylated proteins (PSTPIPs) found in cleavage furrow that are substrates for PEST protein tyrosine phosphatases)

IT Phosphoproteins  
 RL: BOC (Biological occurrence); BPR (Biological process); BSU (Biological study, unclassified); PRP (Properties); BIOL (Biological study); OCCU (Occurrence); PROC (Process)  
 (PSTPIP; tyrosine phosphorylated proteins (PSTPIPs) found in cleavage furrow that are substrates for PEST protein tyrosine phosphatases)

IT Protein motifs  
 (SH3 domain, in PSTPIP; tyrosine phosphorylated proteins (PSTPIPs) found in cleavage furrow that are substrates for PEST protein tyrosine phosphatases)

IT Gene, animal  
 RL: BSU (Biological study, unclassified); PRP (Properties); BIOL (Biological study)  
 (cDNA, for PSTPIP of mouse; tyrosine phosphorylated proteins (PSTPIPs) found in cleavage furrow that are substrates for PEST protein tyrosine phosphatases)

IT Cell division  
 (cytokinesis, PSTPIP in cleavage furrow during; tyrosine phosphorylated proteins (PSTPIPs) found in cleavage furrow that are substrates for PEST protein tyrosine phosphatases)

IT Drug screening  
 (for effectors of PSTPIP-mediated actin polymerization; tyrosine phosphorylated proteins (PSTPIPs) found in cleavage furrow that are substrates for PEST protein tyrosine phosphatases)

IT cDNA sequences  
 (for phosphoprotein PSTPIP of mouse; tyrosine phosphorylated proteins (PSTPIPs) found in cleavage furrow that are substrates for PEST protein tyrosine phosphatases)

IT Antibodies  
 RL: ARG (Analytical reagent use); BUU (Biological use, unclassified); ANST (Analytical study); BIOL (Biological study); USES (Uses)  
 (monoclonal, to PSTPIP; tyrosine phosphorylated proteins (PSTPIPs) found in cleavage furrow that are substrates for PEST protein tyrosine phosphatases)

IT Molecular association  
 (of PSTPIP and PTP PEST, characterization of; tyrosine phosphorylated proteins (PSTPIPs) found in cleavage furrow that are substrates for PEST protein tyrosine phosphatases)

IT Protein motifs  
 (of PSTPIP of mouse; tyrosine phosphorylated proteins (PSTPIPs) found in cleavage furrow that are substrates for PEST protein tyrosine phosphatases)

IT Dephosphorylation, biological  
 (of PSTPIP proteins by PEST phosphatases; tyrosine phosphorylated proteins (PSTPIPs) found in cleavage furrow that are substrates for PEST protein tyrosine phosphatases)

IT Protein sequences  
 (of phosphoprotein PSTPIP of mouse; tyrosine phosphorylated proteins (PSTPIPs) found in cleavage furrow that are substrates for PEST protein tyrosine phosphatases)

IT Antibodies  
 RL: ARG (Analytical reagent use); BUU (Biological use, unclassified); ANST (Analytical study); BIOL (Biological study); USES (Uses)  
 (to PSTPIP; tyrosine phosphorylated proteins (PSTPIPs) found in cleavage furrow that are substrates for PEST protein tyrosine phosphatases)

IT 196523-73-6  
 RL: BOC (Biological occurrence); BSU (Biological study, unclassified); PRP

(Properties); BIOL (Biological study); OCCU (Occurrence)  
 (amino acid sequence, association with actin of; tyrosine phosphorylated  
 proteins (PSTPIPs) found in cleavage furrow that are substrates for  
 PEST protein tyrosine phosphatases)

IT 196717-98-3  
 RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL  
 (Biological study); PROC (Process)  
 (dephosphorylation of PSTPIP by; tyrosine phosphorylated proteins  
 (PSTPIPs) found in cleavage furrow that are substrates for PEST  
 protein tyrosine phosphatases)

IT 211623-46-0  
 RL: BSU (Biological study, unclassified); PRP (Properties); BIOL  
 (Biological study)  
 (nucleotide sequence; tyrosine phosphorylated proteins (PSTPIPs) found  
 in cleavage furrow that are substrates for PEST protein  
 tyrosine phosphatases)

RE.CNT 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD

RE  
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L51 ANSWER 1 OF 11 EMBASE COPYRIGHT 2004 ELSEVIER INC. ALL RIGHTS RESERVED.  
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AN 2003465693 EMBASE  
 TI Molecular Cloning and Characterization of a Novel Dual Specificity  
 Phosphatase, LMW-DSP2, that Lacks the Cdc25 Homology Domain.  
 AU Aoyama K.; Nagata M.; Oshima K.; Matsuda T.; Aoki N.  
 CS .naoki@agr.nagoya-u.ac.jp  
 SO Journal of Biological Chemistry, (20 Jul 2001) 276/29 (27575-27583).  
 Refs: 46  
 ISSN: 0021-9258 CODEN: JBCHA3  
 CY United States  
 DT Journal; Article  
 FS 029 Clinical Biochemistry  
 LA English  
 SL English  
 AB A novel dual specificity phosphatase (DSP) designated LMW-DSP2 was cloned  
 with a combination of reverse transcription-polymerase chain reaction and  
 cDNA library screening strategies. The LMW-DSP2 open reading frame of 194  
 amino acids contained a single DSP catalytic domain but lacked the cdc25  
 homology domain, which is conserved in most known DSPs. Northern blot and  
 reverse transcription-polymerase chain reaction analyses revealed that  
 LMW-DSP2 was specifically expressed in testis. Recombinant LMW-DSP2  
 protein exhibited phosphatase activity toward an artificial low molecular  
 weight substrate paranitrophenyl phosphate, and the activity was inhibited  
 completely by sodium orthovanadate but not sodium fluoride, pyrophosphate,  
 and okadaic acid. The substitution of critical amino acid residues,  
 aspartic acid and cysteine, resulted in a dramatic reduction of  
 phosphatase activity. Transient transfection of LMW-DSP2 in COS7 cells  
 resulted in the expression of a 21-kDa protein, and the phosphatase was  
 shown to be distributed in both the cytosol and the nucleus. LMW-DSP2  
 dephosphorylated and deactivated p38, to a higher extent, and  
 stress-activated protein kinase (SAPK)/c-Jun N-terminal kinase (JNK), but  
 not extracellular signal-regulated kinase 1/2 mitogen-activated protein  
 kinases, in transfected COS7 cells and in vitro. Interestingly, mutation  
 in a conserved docking motif of p38 and SAPK/JNK as well as in a cluster  
 of aspartic acids of LMW-DSP2 did not affect the deactivation of the  
 mitogen-activated protein kinases by LMW-DSP2. Furthermore, the binding  
 between LMW-DSP2 and p38 and SAPK/JNK was also not disrupted by such

mutations. Among the DSPs lacking the cdc25 homology domain, LMW-DSP2 is the first one that dephosphorylates and deactivates p38 and SAPK/JNK.

CT Medical Descriptors:  
 \*molecular cloning  
 \*enzyme specificity  
 \*protein domain  
 \*sequence homology  
 \*nucleotide sequence  
 reverse transcription polymerase chain reaction  
 DNA library  
 genetic screening  
 open reading frame  
 amino acid analysis  
 Northern blotting  
 protein expression  
 testis  
 enzyme activity  
 molecular weight  
 enzyme substrate  
 amino acid substitution  
 genetic transfection  
 enzyme localization  
 cytosol  
 cell nucleus  
 enzyme inactivation  
 protein phosphorylation  
 in vitro study  
 protein motif  
 binding kinetics  
 binding affinity  
 gene mutation  
 nonhuman  
 controlled study  
 animal cell  
 article  
 priority journal  
 Drug Descriptors:  
 \*phosphatase  
 \*dual specificity phosphatase  
 \*protein lmw dsp2  
 \*protein tyrosine phosphatase  
 4 nitrophenyl phosphate  
 vanadate sodium  
 fluoride sodium  
 pyrophosphate  
 okadaic acid  
 aspartic acid  
 cysteine  
 stress activated protein kinase: EC, endogenous compound  
 mitogen activated protein kinase 1: EC, endogenous compound  
 mitogen activated protein kinase 2: EC, endogenous compound  
 unclassified drug

RN (phosphatase) 9013-05-2; (protein tyrosine phosphatase) 79747-53-8, 97162-86-2; (4 nitrophenyl phosphate) 330-13-2; (vanadate sodium) 11105-06-9, 13718-26-8, 13721-39-6; (fluoride sodium) 51668-54-3, 7681-49-4, 79933-27-0; (pyrophosphate) 14000-31-8, 7722-88-5, 7758-16-9; (okadaic acid) 78111-17-8; (aspartic acid) 56-84-8, 6899-03-2; (cysteine) 4371-52-2, 52-89-1, 52-90-4; (stress activated protein kinase) 155215-87-5; (mitogen activated protein kinase 1) 137632-07-6; (mitogen activated protein kinase 2) 137632-08-7

GEN GENBANK AF237619 submitted number; GENBANK M32599 referred number; GENBANK U10871 referred number; GENBANK U81823 referred number; GENBANK Y13439 referred number; GENBANK AB005663 referred number; GENBANK AF135185 referred number; GENBANK AB005664 referred number; GENBANK AB005665 referred number

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 on STN

AN 2003453747 EMBASE

TI Acquisition of a Specific and Potent PTP1B Inhibitor from a Novel Combinatorial Library and Screening Procedure.

AU Shen K.; Keng Y.-F.; Wu L.; Guo X.-L.; Lawrence D.S.; Zhang Z.-Y.

CS D.S. Lawrence, Dept. of Biochemistry, Albert Einstein College of Medicine, Yeshiva University, 1300 Morris Park Ave., Bronx, NY 10461, United States.  
 dlawrenc@aecom.yu.edu

SO Journal of Biological Chemistry, (14 Dec 2001) 276/50 (47311-47319).  
 Refs: 52  
 ISSN: 0021-9258 CODEN: JBCHA3

CY United States  
 DT Journal; Article  
 FS 029 Clinical Biochemistry  
 037 Drug Literature Index

LA English  
 SL English

AB Protein-tyrosine phosphatases (PTPases) form a large family of enzymes that serve as key regulatory components in signal transduction pathways. Defective or inappropriate regulation of PTPase activity leads to aberrant tyrosine phosphorylation, which contributes to the development of many human diseases including cancers and diabetes. For example, recent gene knockout studies in mice identify PTP1B as a promising target for anti-diabetes/obesity drug discovery. Thus, there is intense interest in obtaining specific and potent PTPase inhibitors for biological studies and pharmacological development. However, given the highly conserved nature of the PTPase active site, it is unclear whether selectivity in PTPase inhibition can be achieved. We describe a combinatorial approach that is designed to target both the active site and a unique peripheral site in PTP1B. Compounds that can simultaneously associate with both sites are expected to exhibit enhanced affinity and specificity. We also describe a novel affinity-based high-throughput assay procedure that can be used for PTPase inhibitor screening. The combinatorial library/high-throughput screen protocols furnished a small molecule PTP1B inhibitor that is both potent ( $K(i) = 2.4$  nM) and selective (little or no activity against a panel of phosphatases including Yersinia PTPase, SHP1, SHP2, LAR, HePTP, PTP.alpha., CD45, VHR, MKP3, Cdc25A, Stpl, and PP2C). These results demonstrate that it is possible to acquire potent, yet highly selective inhibitors for individual members of the large PTPase family of enzymes.

CT Medical Descriptors:  
 \*enzyme inhibition  
 \*inhibition kinetics  
 combinatorial library  
 DNA screening  
 protein family  
 regulatory mechanism  
 signal transduction  
 enzyme regulation  
 enzyme phosphorylation  
 cancer risk  
 diabetes mellitus  
 enzyme specificity  
 drug potency  
 enzyme active site  
 binding site  
 binding affinity  
 Yersinia  
 nonhuman  
 article  
 priority journal  
 Drug Descriptors:  
 \*protein tyrosine phosphatase inhibitor: PD, pharmacology  
 \*protein tyrosine phosphatase 1b inhibitor: PD, pharmacology  
 phosphotransferase: EC, endogenous compound  
 protein shp1: EC, endogenous compound  
 protein shp2: EC, endogenous compound  
 protein lar: EC, endogenous compound  
 protein heptp: EC, endogenous compound  
 protein tyrosine phosphatase alpha: EC, endogenous compound  
 CD45 antigen: EC, endogenous compound  
 protein vhr: EC, endogenous compound  
 protein Mkp3: EC, endogenous compound  
 protein tyrosine phosphatase: EC, endogenous compound  
 protein cdc25a: EC, endogenous compound  
 protein stpl: EC, endogenous compound  
 protein pp2c: EC, endogenous compound  
 unclassified drug

RN (phosphotransferase) 9031-09-8, 9031-44-1; (protein tyrosine phosphatase) 79747-53-8, 97162-86-2

L51 ANSWER 3 OF 11 EMBASE COPYRIGHT 2004 ELSEVIER INC. ALL RIGHTS RESERVED.  
 on STN  
 AN 2001438829 EMBASE

TI Structure of protein tyrosine phosphatase 1B  
 in complex with inhibitors bearing two phosphotyrosine mimetics.  
 AU Jia Z.; Ye Q.; Dinaut A.N.; Wang Q.; Waddleton D.; Payette P.;  
 Ramachandran C.; Kennedy B.; Hum G.; Taylor S.D.  
 CS S.D. Taylor, Department of Chemistry, University of Waterloo, Waterloo,  
 Ont. N2L 3G1, Canada. s5taylor@sciborg.uwaterloo.ca  
 SO Journal of Medicinal Chemistry, (20 Dec 2001) 44/26 (4584-4594).  
 Refs: 60  
 ISSN: 0022-2623 CODEN: JMCMAR  
 CY United States  
 DT Journal; Article  
 FS 003 Endocrinology  
 030 Pharmacology  
 037 Drug Literature Index  
 LA English  
 SL English  
 AB Protein tyrosine phosphatases (PTPases) are signal-transducing enzymes that dephosphorylate intracellular proteins that have phosphorylated tyrosine residues. It has been demonstrated that protein tyrosine phosphatase 1B (PTP1B) is an attractive therapeutic target because of its involvement in regulating insulin sensitivity (Elcheby et al. Science 1999, 283, 1544-1548). The identification of a second binding site in PTP1B (Puius et al., Proc. Natl. Acad. Sci. U.S.A. 1997, 94, 13420-13425) suggests a new strategy for inhibitor design, where appropriate compounds may be made to simultaneously occupy both binding sites to gain much higher affinity and selectivity. To test this hypothesis and gain further insights into the structural basis of inhibitor binding, we have determined the crystal structure of PTP1B complexed with two non-peptidyl inhibitors, 4 and 5, both of which contain two aryl difluoromethylenephosphonic acid groups, a nonhydrolyzable phosphate mimetic. The structures were determined and refined to 2.35 and 2.50 Å resolution, respectively. Although one of the inhibitors seems to have satisfied the perceived requirement for dual binding, it did not bind both the active site and the adjacent noncatalytic binding site as expected. The second or distal phosphonate group instead extends into the solvent and makes water-mediated interactions with Arg-47. The selectivity of the more potent of these two inhibitors, as well as four other inhibitors bearing two such phosphate mimetics for PTP1B versus seven other PTPases, was examined. In general, selectivity was modest to good when compared to PTPases Cdc25a, PTPmeg-1, PTP.β., and CD45. However, selectivity was generally poor when compared to other PTPases such as SHP-1, SHP-2, and especially TCPTP, for which almost no selectivity was found. The implications these results have concerning the utility of dual-binding inhibitors are discussed.  
 CT Medical Descriptors:  
 signal transduction  
 drug structure  
 dephosphorylation  
 drug targeting  
 insulin sensitivity  
 drug binding site  
 drug selectivity  
 crystal structure  
 complex formation  
 article  
 Drug Descriptors:  
 \*protein tyrosine phosphatase  
 \*protein tyrosine phosphatase 1B  
 \*protein tyrosine phosphatase inhibitor: AN, drug analysis  
 \*protein tyrosine phosphatase inhibitor: DV, drug development  
 \*protein tyrosine phosphatase inhibitor: PD, pharmacology  
 phosphotyrosine  
 unclassified drug  
 RN (protein tyrosine phosphatase) 79747-53-8,  
 97162-86-2; (phosphotyrosine) 21820-51-9  
 L51 ANSWER 4 OF 11 EMBASE COPYRIGHT 2004 ELSEVIER INC. ALL RIGHTS RESERVED.  
 on STN  
 AN 2001176680 EMBASE  
 TI Phospho-azatyrosine, a less effective protein-tyrosine phosphatase substrate than phosphotyrosine.  
 AU Burke T.R. Jr.; Yao Z.-J.; Ye B.; Miyoshi K.; Otaka A.; Wu L.; Zhang Z.-Y.  
 CS T.R. Burke Jr., Division of Basic Sciences, National Cancer Institute,  
 National Institutes of Health, Boyles Street, Frederick, MD 21702-1201,  
 United States. tburke@helix.nih.gov  
 SO Bioorganic and Medicinal Chemistry Letters, (21 May 2001) 11/10

(1265-1268).  
 Refs: 21  
 ISSN: 0960-894X CODEN: BMCL8.  
 PUI S 0960-894X(01)00197-4  
 CY United Kingdom  
 DT Journal; Article  
 FS 030 Pharmacology  
 037 Drug Literature Index  
 LA English  
 SL English  
 AB Azatyrosine (AzaTyr, 4) is a natural product isolated from *Streptomyces chibensis*, whose structure is characterized by a nitrogen atom in the aryl ring of a tyrosyl residue. This seemingly minor modification to the tyrosyl residue results in profound physiological effects, as AzaTyr has been shown to promote permanent reversion of ras-dependent transformed cells to the normal phenotype in culture and to inhibit chemical induction of carcinogenesis in transgenic mice bearing oncogenic human ras. The mechanisms underlying these effects are not known, however ras-pathways involve an intricate balance between both protein-tyrosine kinases (PTKs) and protein-tyrosine phosphatases (PTPs). The present study was undertaken to examine the general utility of AzaTyr as a structural motif for PTP inhibitor design by examining the phospho-azatyrosine (pAzaTyr)-containing peptide Ac-Asp-Ala-Asp-Glu-pAzaTyr-Leu-amide (8) in a PTP1 enzyme system. Kinetic analysis indicated that 8 binds with a  $K_m$  value of 210  $\mu\text{M}$  and a catalytic turnover rate,  $k_{\text{cat}}$  of 52  $\text{s}^{-1}$ . This represents a greater than 50-fold reduction in binding affinity relative to the parent phosphotyrosine-containing peptide, indicating that the aryl nitrogen adversely affects binding affinity. The much lower PTP affinity of the pAzaTyr-containing peptide reduces the potential utility of the AzaTyr pharmacophore for PTP inhibitor design. These results are discussed from the point of view that incorporation of AzaTyr residues into proteins could result in perturbation of protein-tyrosine phosphorylation/dephosphorylation cascades that control signal transduction processes, including ras-dependent pathways.  
 CT Medical Descriptors:  
 drug design  
 peptide analysis  
 kinetics  
 protein binding  
 catalysis  
 binding affinity  
 enzyme binding  
 drug utilization  
 pharmacophore  
 phosphorylation  
 dephosphorylation  
 signal transduction  
 oncogene ras  
 drug structure  
 enzyme inhibition  
 cell culture  
 article  
 Drug Descriptors:  
 \*phosphoazatyrosine: AN, drug analysis  
 \*phosphoazatyrosine: CM, drug comparison  
 \*phosphoazatyrosine: PR, pharmaceutics  
 \*phosphoazatyrosine: PD, pharmacology  
 \*tyrosine derivative: AN, drug analysis  
 \*tyrosine derivative: CM, drug comparison  
 \*tyrosine derivative: PR, pharmaceutics  
 \*tyrosine derivative: PD, pharmacology  
 \*enzyme inhibitor: AN, drug analysis  
 \*enzyme inhibitor: CM, drug comparison  
 \*enzyme inhibitor: PR, pharmaceutics  
 \*enzyme inhibitor: PD, pharmacology  
 \*phosphotyrosine: CM, drug comparison  
 \*phosphotyrosine: PD, pharmacology  
 protein tyrosine phosphatase 1: EC, endogenous compound  
 protein tyrosine phosphatase: EC, endogenous compound  
 acetylaspartylalanylasparylglutamylphosphrylazatyrosineleucine  
 amide: AN, drug analysis  
 acetylaspartylalanylasparylglutamylphosphrylazatyrosineleucine amide: DV,  
 drug development  
 acetylaspartylalanylasparylglutamylphosphrylazatyrosineleucine amide: PR,  
 pharmaceutics

acetylaspartylalanylasparylglutamylphosphorylazatyrosineleucine amide: PD,  
 pharmacology  
 nitrogen  
 unclassified drug  
 RN (tyrosine derivative) 42406-77-9; (phosphotyrosine) 21820-51-9; (protein tyrosine phosphatase) 79747-53-8, 97162-86-2; (nitrogen) 7727-37-9

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 on STN

AN 2000362285 EMBASE

TI Protein tyrosine phosphatases (PTPS) as drug  
 targets: Inhibitors of PTP-1B for the treatment of diabetes.

AU Hundahl Moller N.P.; Iversen L.F.; Andersen H.S.; McCormack J.G.

CS N.P. Hundahl Moller, Signal Transduction, Target Cell Biology, Novo Alle, DK-2880 Bagsvaerd, Denmark. nphm@novo.dk

SO Current Opinion in Drug Discovery and Development, (2000) 3/5 (527-540).  
 Refs: 115  
 ISSN: 1367-6733 CODEN: CODDF

CY United Kingdom

DT Journal; General Review

FS 030 Pharmacology  
 037 Drug Literature Index

LA English

SL English

AB The phosphorylation of key proteins on tyrosine residues is an important part of many different intracellular signaling cascade mechanisms triggered by hormones and other agents. The deactivation of such signaling processes is catalyzed by protein tyrosine phosphatases (PTPs), and therefore inhibition of these enzymes is being explored in different indications as a means whereby signaling may be prolonged or even initiated in the absence of the triggering agent. In the case of the signaling cascade initiated by the activation of the insulin receptor, an important gene knockout study in mice has identified PTP-1B as a potential target 'for antidiabetes therapy, and has thus made it a 'focus of attention' for several groups. Recent advances in the structure-based design of potent and selective inhibitors of this enzyme are described, as well as some preliminary data for such inhibitors in animal models which, together with more recently published data from further studies on PTP-1B knockout mice and from antisense studies, illustrate the potential of this approach for the treatment of both Type I and Type II diabetes.

CT Medical Descriptors:  
 \*diabetes mellitus: DT, drug therapy  
 drug design  
 drug screening  
 drug receptor binding  
 signal transduction  
 enzyme inhibition  
 insulin sensitivity  
 enzyme phosphorylation  
 protein domain  
 knockout mouse  
 glucose homeostasis  
 human  
 nonhuman  
 review  
 Drug Descriptors:  
 \*protein tyrosine phosphatase: EC, endogenous compound  
 \*protein tyrosine phosphatase inhibitor: AN, drug analysis  
 \*protein tyrosine phosphatase inhibitor: DV, drug development  
 \*protein tyrosine phosphatase inhibitor: DT, drug therapy  
 \*protein tyrosine phosphatase inhibitor: PD, pharmacology  
 glucose  
 isoenzyme: EC, endogenous compound  
 protein tyrosine phosphatase 1b: EC, endogenous compound  
 insulin receptor: EC, endogenous compound  
 insulin receptor kinase: EC, endogenous compound  
 CD45 antigen: EC, endogenous compound  
 acid phosphatase: EC, endogenous compound  
 antisense oligonucleotide  
 leptin: EC, endogenous compound  
 insulin: EC, endogenous compound  
 triacylglycerol: EC, endogenous compound  
 naphthalene derivative: AN, drug analysis  
 naphthalene derivative: DV, drug development  
 naphthalene derivative: PD, pharmacology

vanadium derivative: AN, drug analysis

vanadium derivative: DT, drug therapy

vanadium derivative: PD, pharmacology

2 (oxalylamino)benzoic acid: AN, drug analysis

2 (oxalylamino)benzoic acid: DV, drug development

2 (oxalylamino)benzoic acid: PD, pharmacology

unclassified drug

RN (protein tyrosine phosphatase) 79747-53-8,  
97162-86-2; (glucose) 50-99-7, 84778-64-3; (acid phosphatase) 9001-77-8,  
9025-88-1; (insulin) 9004-10-8

CO Novo Nordisk; Merck Frosst; Pharmacia Upjohn; Home Products

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AN 2000053341 EMBASE

TI Structure-based discovery of small molecule inhibitors targeted to  
protein tyrosine phosphatase 1B.

AU Sarmiento M.; Wu L.; Keng Y.-F.; Song L.; Luo Z.; Huang Z.; Wu G.-Z.; Yuan  
A.K.; Zhang Z.-Y.

CS Z.-Y. Zhang, Department of Molecular Pharmacology, Albert Einstein College  
of Medicine, 1300 Morris Park Avenue, Bronx, NY 10461, United States.

zhang@aecon.yu.edu

SO Journal of Medicinal Chemistry, (27 Jan 2000) 43/2 (146-155).

Refs: 64

ISSN: 0022-2623 CODEN: JMCMAR

CY United States

DT Journal; Article

FS 029 Clinical Biochemistry

030 Pharmacology

037 Drug Literature Index

LA English

SL English

AB Protein tyrosine phosphatases (PTPases) are involved in the control of tyrosine phosphorylation levels in the cell and are believed to be crucial for the regulation of a multitude of cellular functions. A detailed understanding of the role played by PTPases in various signaling pathways has not yet been achieved, and potent and selective PTPase inhibitors are essential in the quest to determine the functionality of individual PTPases. Using the DOCK methodology, we have carried out a structure-based, computer-assisted search of an available chemical database in order to identify low molecular weight, nonpeptidic PTP1B inhibitors. We have identified several organic molecules that not only possess inhibitory activity against PTP1B but which also display significant selectivity for PTP1B. This indicates that although structural features important for pTyr recognition are conserved among different PTPases, it is possible to generate selective inhibitors targeted primarily to the catalytic site. Kinetic analysis and molecular modeling experiments suggest that the PTP1B active site possesses significant plasticity such that substituted and extended aromatic systems can be accommodated. The newly identified molecules provide a molecular framework upon which therapeutically useful compounds can ultimately be based, and systematic optimization of these lead compounds is likely to further enhance their potency and selectivity.

CT Medical Descriptors:

\*drug targeting

protein phosphorylation

cell function

signal transduction

structure activity relation

molecular interaction

article

Drug Descriptors:

\*protein tyrosine phosphatase inhibitor: AN, drug analysis

\*protein tyrosine phosphatase inhibitor: DV, drug development

\*protein tyrosine phosphatase inhibitor: PD, pharmacology

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AN 2000053013 EMBASE

TI Synthesis and biological evaluation of a targeted library of protein  
phosphatase inhibitors.

AU Wipf P.; Asian D.C.; Luci D.K.; Southwick E.C.; Lazo J.S.

CS P. Wipf, Department of Chemistry, University of Pittsburgh, Pittsburgh, PA  
15260, United States

SO Biotechnology and Bioengineering, (2000) 71/1 (58-70).

Refs: 34

ISSN: 0006-3592 CODEN: BIBIAU  
 CY United States  
 DT Journal; Article  
 FS 030 Pharmacology  
 037 Drug Literature Index  
 LA English  
 SL English  
 AB Phosphorylation of serine, threonine, and tyrosine controls fundamental mammalian cell events and is achieved by kinases which, in turn, are in dynamic relationship with phosphatases. Few selective inhibitors of protein tyrosine and dual specificity phosphatases are readily available. Based on SAR studies of naturally occurring phosphatase inhibitors and following up on previously published research, we have designed a new pharmacophore model V and synthesized a new library of functional analogues of V. All synthetic steps were carried out and optimized employing combinatorial chemistry methods on Wang resin. All compounds were tested in vitro for their ability to inhibit recombinant human protein tyrosine (PTP1B) and dual-specificity (Cdc25B2 and VHR) phosphatases. Three of the approximately 70 compounds in our library inhibited Cdc25B2 by 50% at 375-490  $\mu$ M. No compounds inhibited PTP1B, and only one blocked VHR. Cell-culture studies revealed no toxicity to human breast cancer cells with two of the phosphatase inhibitors. (C) 2000 John Wiley and Sons, Inc.  
 CT Medical Descriptors:  
 \*drug synthesis  
 protein phosphorylation  
 structure activity relation  
 in vitro study  
 pharmacophore  
 enzyme inhibition  
 enzyme activity  
 breast cancer  
 cell proliferation  
 human  
 controlled study  
 human cell  
 article  
 Drug Descriptors:  
 \*phosphoprotein phosphatase inhibitor: AN, drug analysis  
 \*phosphoprotein phosphatase inhibitor: DV, drug development  
 \*phosphoprotein phosphatase inhibitor: PD, pharmacology  
 protein tyrosine phosphatase  
 phosphoprotein phosphatase  
 resin  
 cyanoginosin LR  
 okadaic acid  
 calyculin A  
 sulfonamide  
 amine  
 amide  
 lysine derivative  
 (protein tyrosine phosphatase) 79747-53-8,  
 97162-86-2; (phosphoprotein phosphatase) 9025-75-6; (cyanoginosin LR)  
 101043-37-2; (okadaic acid) 78111-17-8; (calyculin A) 101932-71-2; (amide)  
 17655-31-1  
 L51 ANSWER 8 OF 11 EMBASE COPYRIGHT 2004 ELSEVIER INC. ALL RIGHTS RESERVED.  
 on STN  
 AN 1998394517 EMBASE  
 TI Src Homology-2 Domains: Structure, mechanisms, and drug discovery.  
 AU Sawyer T.K.  
 CS T.K. Sawyer, ARIAD Pharmaceuticals, Inc., 26 Landsdowne St., Cambridge, MA 02139, United States. tomi.sawyer@ariad.com  
 SO Biopolymers - Peptide Science Section, (1998) 47/3 (243-261).  
 Refs: 44  
 ISSN: 0006-3525 CODEN: BPSSFT  
 CY United States  
 DT Journal; General Review  
 FS 029 Clinical Biochemistry  
 037 Drug Literature Index  
 LA English  
 SL English  
 AB Src homology-2 (SH2) domains and their associated catalytic or noncatalytic proteins constitute critical signal transduction targets for drug discovery. Such SH2 proteins are found in the regulation of a number of cellular processes, including growth, mitogenesis, motility,

metabolism, immune response, and gene transcription. From the relationship of tyrosine phosphorylation and intracellular regulation by protein-tyrosine kinases (PTKs) and protein-tyrosine phosphatases (PTPs), the dynamic and reversible binding interactions of SH2 domain containing proteins with their cognate phosphotyrosine (pTyr) containing proteins provide a third dimensionality to the orchestration of signal transduction pathways that exist as a result of pTyr formation, degradation, and molecular recognition events. This review highlights several key research achievements impacting our current understanding of SH2 structure, mechanisms, and drug discovery that underlie the role(s) of SH2 domains in signal transduction processes, cellular functions, and disease states.

CT Medical Descriptors:

- \*protein structure
- \*sequence homology
  - \*drug screening
- \*signal transduction
- catalysis
- cell growth
- mitogenesis
- cell motility
- cell metabolism
- immune response
- genetic transcription
  - protein phosphorylation
- protein protein interaction
- molecular recognition
- drug targeting
- drug design
- structure activity relation
- human
- nonhuman
- review

Drug Descriptors:

- \*protein tyrosine kinase: EC, endogenous compound
- \*protein kinase p60: EC, endogenous compound
  - \*protein tyrosine phosphatase: EC, endogenous compound
- phosphotyrosine: EC, endogenous compound
  - phosphopeptide
- peptide library
  - protein tyrosine kinase inhibitor: AN, drug analysis
  - protein tyrosine kinase inhibitor: DV, drug development
  - protein kinase inhibitor: AN, drug analysis
  - protein kinase inhibitor: DV, drug development
  - protein tyrosine phosphatase inhibitor: AN, drug analysis
  - protein tyrosine phosphatase inhibitor: DV, drug development
  - nonapeptide: AN, drug analysis
  - nonapeptide: DV, drug development
- prodrug

RN (protein tyrosine kinase) 80449-02-1; (protein tyrosine phosphatase) 79747-53-8, 97162-86-2; (phosphotyrosine) 21820-51-9

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AN 1998394516 EMBASE

TI Protein-tyrosine phosphatases: Structure, mechanism, and inhibitor discovery.

AU Burke T.R. Jr.; Zhang Z.-Y.

CS T.R. Burke Jr., Building 37, National Institutes of Health, Bethesda, MD 20892, United States

SO Biopolymers - Peptide Science Section, (1998) 47/3 (225-241).

Refs: 190

ISSN: 0006-3525 CODEN: BPSSFT

CY United States

DT Journal; General Review

FS 029 Clinical Biochemistry

037 Drug Literature Index

LA English

SL English

AB Protein-tyrosine kinases (PTKs) and their associated signaling pathways are crucial for the regulation of numerous cell functions including growth, mitogenesis, motility, cell-cell interactions, metabolism, gene transcription, and the immune response. Since tyrosine phosphorylation is reversible and dynamic in vivo, the phosphorylation states of proteins are governed by the opposing actions of PTKs and protein-tyrosine phosphatases (PTPs). In this light, both PTKs

and PTPs play equally important roles in signal transduction in eukaryotic cells, and comprehension of mechanisms behind the reversible pTyr-dependent modulation of protein function and cell physiology must necessarily encompass the characterization of PTPs as well as PTKs. In spite of the large number of PTPs identified to date and the emerging role played by PTPs in disease, a detailed understanding of the role played by PTPs in signaling pathways has been hampered by the absence of PTP-specific agents. Such PTP-specific inhibitors could potentially serve as useful tools in determining the physiological significance of protein tyrosine phosphorylation in complex cellular signal transduction pathways and may constitute valuable therapeutics in the treatment of several human diseases. The goal of this review is therefore to summarize currently understandings of PTP structure and mechanism of catalysis and the relationship of these to PTP inhibitor development. The review is organized such that enzyme structure is covered first, followed by mechanisms of catalysis then PTP inhibitor development. In discussing PTP inhibitor development, nonspecific inhibitors and those obtained by screening methods are initially presented with the focus then shifting to inhibitors that utilize a more structure-based rationale.

CT

Medical Descriptors:

- \*enzyme structure
- \*enzyme mechanism

- \*protein phosphorylation
- \*drug screening

signal transduction

catalysis

cell growth

mitogenesis

cell motility

cell interaction

cell metabolism

genetic transcription

immune response

osteoclast

human

nonhuman

review

Drug Descriptors:

- \*protein tyrosine phosphatase: EC, endogenous compound

- \*protein tyrosine phosphatase inhibitor: DV, drug development

natural product

apomorphine

acid phosphatase prostate isoenzyme

alkaline phosphatase bone isoenzyme

phosphonic acid derivative: DV, drug development

alendronic acid: DV, drug development

peptide library

RN (protein tyrosine phosphatase) 79747-53-8,  
97162-86-2; (apomorphine) 314-19-2, 58-00-4; (alendronic acid) 66376-36-1

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AN 97331069 EMBASE

DN 1997331069

TI A combinatorial approach to identifying protein tyrosine phosphatase substrates from a phosphotyrosine peptide library.

AU Cheung Y.W.; Abell C.; Balasubramanian S.

CS S. Balasubramanian, University Chemical Laboratory, Lensfield Road, Cambridge CB2 1EW, United Kingdom

SO Journal of the American Chemical Society, (1997) 119/40 (9568-9569).

Refs: 25

ISSN: 0002-7863 CODEN: JACSAT

CY United States

DT Journal; Article

FS 029 Clinical Biochemistry

LA English

CT Medical Descriptors:

- \*enzyme specificity

- \*protein phosphorylation

article

methodology

- screening

sequence analysis

Drug Descriptors:

- \*phosphotyrosine

- \*protein tyrosine phosphatase

RN (phosphotyrosine) 21820-51-9; (protein tyrosine phosphatase) 79747-53-8, 97162-86-2  
 L51 ANSWER 11 OF 11 EMBASE COPYRIGHT 2004 ELSEVIER INC. ALL RIGHTS RESERVED.  
 on STN  
 AN 95359852 EMBASE  
 DN 1995359852  
 TI Synthetic Tyr-phospho and non-hydrolyzable phosphonopeptides as PTKs and TC-PTP inhibitors.  
 AU Ruzza P.; Deana A.D.; Calderan A.; Pavanetto M.; Cesaro L.; Pinna L.A.; Borin G.  
 CS Centro di Studio sui Biopolimeri, CNR, Dipart. di Chimica Organica, Universita di Padova, Via Marzolo 1,35131 Padova, Italy  
 SO International Journal of Peptide and Protein Research, (1995) 46/6 (535-546).  
 ISSN: 0367-8377 CODEN: IJPPC3  
 CY Denmark  
 DT Journal; Article  
 FS 029 Clinical Biochemistry  
 037 Drug Literature Index  
 LA English  
 SL English  
 AB Tyrosine-specific protein kinases and phosphatases are important signal transducing enzymes in normal cellular growth and differentiation and have been implicated in the etiology of a number of human neoplastic processes. In order to develop agents which inhibit the function of these two classes of enzymes by interfering with the binding of their substrates, we synthesized analogs derived from the peptide EDNEYTA. This sequence reproduces the main autophosphorylation site of Src tyrosine kinases. In this work we report the synthesis, by classical solution methods, of the phosphotyrosyl peptide EDNEYpTA as well as of three analogs in which the phosphotyrosine is replaced by a phosphotyrosine and by two unnatural, non-hydrolyzable amino acids 4-phosphonomethyl-L-phenylalanine and 4-phosphono-L-phenylalanine. The Src peptide and its derivatives were tested as inhibitors of three non-receptor tyrosine kinases (Lyn, belonging to the Src family, CSK and PTK-IIB) and a non-receptor protein tyrosine phosphatase obtained from human T-cell (TC-PTP). The biomimetic analogues, which do not significantly affect the activity of CSK, PTK-IIB and TC-PTP, act as efficient inhibitors on Lyn, influencing both the exogenous phosphorylation and, especially, its autophosphorylation. In particular, the Pphe derivative may provide a basis for the design of a class of inhibitors specific for Lyn and possibly Src tyrosine kinases, capable of being used in vivo and in vitro conditions.  
 CT Medical Descriptors:  
 \*enzyme inhibition  
 article  
 autophosphorylation  
 circular dichroism  
 controlled study  
 drug design  
 human  
 human cell  
 peptide synthesis  
 protein phosphorylation  
 protein structure  
 structure activity relation  
 t lymphocyte  
 Drug Descriptors:  
 \*enzyme inhibitor: AN, drug analysis  
 \*enzyme inhibitor: CM, drug comparison  
 \*enzyme inhibitor: DV, drug development  
 \*protein tyrosine kinase: PR, pharmaceutics  
 \*protein tyrosine phosphatase: EC, endogenous compound  
 \*synthetic peptide: AN, drug analysis  
 \*synthetic peptide: CM, drug comparison  
 \*synthetic peptide: DV, drug development  
 RN (protein tyrosine kinase) 80449-02-1; (protein tyrosine phosphatase) 79747-53-8, 97162-86-2

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L60 ANSWER 1 OF 3 WPIX COPYRIGHT 2004 THE THOMSON CORP on STN  
 AN 2001-235905 [25] WPIX  
 DNC C2001-070998  
 TI Identifying a combination of a dephosphorylating enzyme and a  
 phosphorylated protein that forms a complex involved in the control of  
 cell regulation comprises oxidatively deactivating the enzyme.  
 DC B04 D16  
 IN BOEHMER, F; HERRLICH, P  
 PA (GESL) FORSCHUNGSZENTRUM KARLSRUHE GMBH  
 CYC 20  
 PI DE 10035472 A1 20010315 (200125)\* 25 C12N009-14  
 WO 2001020021 A2 20010322 (200125) GE C12Q001-42  
 RW: AT BE CH CY DE DK ES FI FR GB GR IE IT LU MC NL PT SE  
 W: JP US  
 ADT DE 10035472 A1 DE 2000-10035472 20000721; WO 2001020021 A2 WO 2000-EP7455  
 20000802  
 PRAI DE 1999-19944069 19990914  
 IC ICM C12N009-14; C12Q001-42  
 ICS A61K038-17; A61K038-45; A61K038-46; C07K014-435; C07K014-71;  
 C12N009-12; C12N009-16; C12N013-00; C12Q001-48; G01N033-68  
 AB DE 10035472 A UPAB: 20010508  
 NOVELTY - Method (M) for identifying a combination of a  
 dephosphorylating enzyme and a phosphorylated protein that forms a  
 complex involved in the control of cell regulation, in new  
 DETAILED DESCRIPTION - Method (M) for identifying a combination of a  
 dephosphorylating enzyme and a phosphorylated protein that forms a  
 complex involved in the control of cell regulation comprises:  
 (a) providing a system comprising at least one  
 dephosphorylating enzyme and at least one phosphorylated protein;  
 (b) deactivating the enzyme(s) by oxidation;  
 (c) isolating any complex not undergoing catalytic conversion; and  
 (d) identifying the components of the complex.  
 INDEPENDENT CLAIMS are also included for the following:  
 (1) a complex (I) comprising an oxidatively deactivated  
 dephosphorylating enzyme and a phosphorylated protein;  
 (2) the dephosphorylating enzyme (II) contained in (I);  
 (3) the phosphorylated protein (III) contained in (I);  
 (4) a substrate of (III) that forms a complex with (II);  
 (5) a method (M1) for deactivating a dephosphorylating  
 enzyme, comprising exposing the enzyme, or cells containing it, to  
 radiation, oxidizing agents and/or alkylating agents;  
 (6) dephosphorylated (sic) enzymes produced by (M1);  
 (7) a screening (M2) assay for effectors of (I), (II) or (III),  
 comprising:  
 (a) either incubating (I), (II) or (III) with at least one test  
 substance or irradiating (I), (II) or (III);  
 (b) measuring the specific activity of (II) and/or degree of  
 phosphorylation of (III);  
 (c) repeating step (b) in the absence of the test substance(s) or  
 without irradiation; and  
 (d) comparing the results from steps (b) and (c);

(8) effectors identified by (M2).

USE - Enzyme-protein (especially phosphatase-kinase) pairs identified by the method are useful as targets in screening assays and drug development programs aimed at finding agents capable of modulating the control mechanisms of cell regulation, signal transduction, cell proliferation and/or cell differentiation, especially agents for treating neurodegenerative diseases, diabetes, atherosclerosis or cancer.

Dwg.0/12

FS CPI  
FA AB; DCN  
MC CPI: B04-K01; B04-L04; B04-L05; B11-C08E; B11-C09; B12-K04E;  
D05-H09

TECH UPTX: 20010508  
TECHNOLOGY FOCUS - BIOLOGY - Preferred Method: The enzyme is a protein tyrosine phosphatase and the protein is a protein tyrosine kinase, especially a cell surface receptor with tyrosine kinase activity. The system of step (a), of the method (M), is an artificial system or a natural system, preferably a living cell, especially a mammalian cell. The enzyme is reversibly deactivated so that it binds to the protein without catalytically converting it, preferably by oxidizing an amino acid at its catalytic activity center, either by exposure to radiation, especially ultraviolet radiation with a wavelength of 335, 312 or 200-280 nm, or by treatment with an oxidizing agent, especially hydrogen peroxide, or an alkylating agent.

L60 ANSWER 2 OF 3 WPIX COPYRIGHT 2004 THE THOMSON CORP on STN  
AN 1999-327025 [27] WPIX  
DNN N1999-245288 DNC C1999-096772  
TI Identifying modulators agents that modulate leptin activity.  
DC B04 D16 S03  
IN FRIEDMAN, J M; LI, C  
PA (UYRQ) UNIV ROCKEFELLER  
CYC 21  
PI WO 9923493 A1 19990514 (199927)\* EN 86 G01N033-68  
RW: AT BE CH CY DE DK ES FI FR GB GR IE IT LU MC NL PT SE  
W: CA JP MX  
ADT WO 9923493 A1 WO 1998-US22797 19981027  
PRAI US 1998-178691 19981026; US 1997-961809 19971031  
IC ICM G01N033-68  
ICS G01N033-573; G01N033-74  
AB WO 9923493 A UPAB: 19990714  
NOVELTY - A method for identifying agents that modulate leptin activity, is new.

DETAILED DESCRIPTION - The method of identifying a modulator of binding of a phosphorylated leptin receptor with tyrosine phosphatase 1D (PTP-1D) comprises:

(a) contacting a tyrosine-985 phosphorylated leptin receptor or its phosphorylated fragment with protein tyrosine phosphatase 1D (PTP-1D) or its fragment in the presence and absence of a candidate agent under conditions in which in the absence of the agent the binding of the phosphorylated leptin receptor or fragment with PTP-1D or its fragment can be detected; and

(b) detecting the binding of the phosphorylated leptin receptor and PTP 1d;

where an increase in binding detected in the presence of the agent, indicates that the agent enhances binding, and a decrease in binding in the presence of the agent indicates that the agent is a binding inhibitor.

INDEPENDENT CLAIMS are also included for the following:

(1) identifying modulators of phosphorylated leptin receptor-dependent PTP-1D phosphorylation, optionally in situ;  
(2) identifying modulators of leptin-dependent PTP-1D dephosphorylation of JAK2 kinase in situ;  
(3) identifying inhibitors of leptin-dependent PTP-1D phosphorylation in situ; and  
(4) identifying drugs useful in a weight loss diet regimen.

ACTIVITY - Anorectic.

MECHANISM OF ACTION - Enzyme Inhibitor.

USE - Modulators of tyrosine-985-phosphorylated leptin receptor-dependent PTP-1D phosphorylation are useful as drugs in weight loss diet regimens. The drugs identified can regulate adiposity and fat content of animals, particularly in mammals. Disorders that can be treated by PTP-1D modulators include obesity and its associated diseases, e.g. hypertension, heart disease and type II diabetes, and weight loss associated with cancer and AIDS. Additionally the agents identified may be useful in agriculture where body weight of domestic animals can be

modulated.  
 FS CPI EPI  
 FA AB; DCN  
 MC CPI: B04-K01; B11-C08; B12-K04A; D05-H09; D05-H10  
 EPI: S03-E14H; S03-E14H4  
 TECH UPTX: 19990714

TECHNOLOGY FOCUS - BIOTECHNOLOGY - Preferred Methods: The phosphorylated leptin receptor or its phosphorylated fragment is bound to a solid support. The phosphorylated fragment is part of a fusion protein, i.e. fused to glutathione-S-transferase or green fluorescent protein. PTP-1D or its fragment is labeled.

Modulators of tyrosine-985 phosphorylated leptin receptor dependent PTP-1D phosphorylation can be identified by contacting the receptor with PTP-1D and JAK2 kinase in the presence and absence of an agent. Absence of the agent allows phosphorylation of PTP-1D. The amount of PTP-1D phosphorylation is measured in the presence and absence of the agent. Potential modulators are then contacted with PTP-1D and JAK2 in the absence of a phosphorylated receptor. When no significant change in phosphorylation is determined in the presence of the potential modulator relative to that in the absence of the potential modulator, the potential modulator is a modulator of the leptin-dependent phosphorylation of PTP-1D.

The method of (2) comprises contacting a cell with leptin in the presence or absence of an agent under conditions in which in the absence of the agent leptin induces the phosphorylation of PTP-1D, where the cell comprises PTP-1D, JAK2, and a tyrosine-985 leptin receptor. The amount of PTP-1D phosphorylation is then measured, where an increase or decrease in phosphorylation of PTP-1D is determined in the presence of the agent relative to in the absence of the agent. This method uses cells transfected with vectors encoding PTP-1D, JAK2 and a leptin receptor containing tyrosine-985. The method of (2) further comprises contacting a second cell with leptin and the potential modulator under the conditions described above except where the leptin cannot induce phosphorylation of PTP-1D and where the second cell is transfected with vectors encoding PTP-1D, JAK2 and a leptin receptor that does not contain a tyrosine-985. A modulator is identified when there is no significant change in phosphorylation in the presence of the potential modulator relative to in the absence of the potential modulator. The leptin receptor that does not contain tyrosine-985 is Ob-Ra or Ob-Rb containing a phenylalanine-985. The modulator can enhance or inhibit the leptin receptor-dependent phosphorylation of PTP-1D.

An inhibitor of leptin-dependent PTP-1D phosphorylation can be identified in a similar manner, where in the absence of the agent, leptin induces the expression of a reporter gene operably under the control of a promoter containing a binding site for activated Stat3. Modulators of leptin-dependent PTP-1D dephosphorylation of JAK2 kinase can also be identified in situ in a similar manner.

The modulators identified can be administered to test animals. Modulators that causes the test animal to lose weight relative to a control animal (receiving multiple doses of a placebo) are selected as a drug useful in weight loss diet regimens.

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 AN 1997-424288 [39] WPIX  
 DNC C1997-135768  
 TI Protein tyrosine phosphatase src homology domain binding peptide - corresponds to the phosphatase binding site in EPO receptor, used to prolong the effect of EPO and to identify other phosphatase(s).  
 DC B04 D16  
 IN KLINGMUELLER, U; LODISH, H F; MICHNICK, S  
 PA (WHED) WHITEHEAD INST BIOMEDICAL RES  
 CYC 1  
 PI US 5659012 A 19970819 (199739)\* 14 A61K038-04  
 ADT US 5659012 A US 1995-402006 19950310  
 PRAI US 1995-402006 19950310  
 IC ICM A61K038-04  
 AB US 5659012 A UPAB: 19990525  
 A peptide (A) which binds to the src homology-2 domain of protein tyrosine phosphatase (PTP) SH-PTP1 is new. TPPHLKYLYVVS  
 (A) Also claimed are derivatives of (A) having at least one amino acid modified by substitution with a fluoroether, methyl ether or thioether group.  
 USE - (A) represents the site in erythropoietin receptor (EPO-R) to which SH-PTP1 binds, resulting in activation of phosphatase activity, dephosphorylation of the receptor and of JAK2 kinase, so that the

EPO proliferative signal is ended. (A) can be used as an affinity reagent to identify other phosphatases that bind to EPO-R, also therapeutically to prolong the effect of EPO, e.g. where this is being used to stimulate haemoglobin synthesis or to treat anaemia (associated with renal failure, chronic disease, HIV infection, blood loss or cancer).

Dwg. 0/4

FS CPI  
FA AB; DCN  
MC CPI: B04-C01C; B11-C08E3; B12-K04; B14-H01; D05-H17A4

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